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6 Insights from Pb and O isotopes into along-arc variations in subduction inputs and
7 crustal assimilation for volcanic rocks in Java, Sunda arc, Indonesia
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Abstract

New Pb isotope data are presented for Gede Volcanic Complex, Salak and Galunggung volcanoes in West Java, Merbabu and Merapi volcanoes in Central Java and Ijen Volcanic Complex in East Java of the Sunda arc, Indonesia. New O isotope data for Merbabu and new geochemical and radiogenic isotope data (Sr-Nd-Hf-Pb) for three West Javanese, upper crustal, Tertiary sedimentary rocks are also presented. The data are combined with published geochemical and isotopic data to constrain the relative importance of crustal assimilation and subducted input of crustal material in petrogenesis in Java. Also discussed are the significance of limestone assimilation in controlling the geochemical and isotopic characteristics of erupted Javanese rocks and the geochemical impact upon central and eastern Javanese arc rocks due to the subduction of Roo Rise between 105-109°E. The negative correlation between Pb isotopes and SiO₂, combined with mantle-like $\delta^{18}\text{O}$ values in Gede Volcanic Complex rocks, West Java, are most likely explained by assimilation of more isotopically-primitive arc rocks and/or ophiolitic crust known to outcrop in West Java. The negative Pb isotope-SiO₂ trend cannot be explained by assimilation of the known compositions of the upper crustal rocks. A peak in $\delta^{18}\text{O}$ whole-rock and mineral values in Central Javanese volcanic rocks (Merbabu and Merapi) combined with along-arc trends in Sr isotope ratios suggest that a different or additional crustal assimilant exerts control on the isotopic composition of Central Javanese volcanic rocks. This assimilant (likely carbonate material) is characterised by high $\delta^{18}\text{O}$ and high Sr isotope ratio but is not particularly elevated in its Pb isotopic ratio. Once the effects of crustal assimilation are accounted for, strong East to West Java regional variations in Ba concentration, Ba/Hf ratio and Pb isotopic composition are evident. These differences are attributed to heterogeneity in the subducted source input component along the island: a more radiogenic Pb isotopic, lower Ba/Hf component (detrital-rich subducted sediment) in West Java and a less radiogenic Pb isotopic, high Ba/Hf component (attributed to a greater AOC/sediment fluid component and/or dominance of pelagic, clay-rich subducted sediment) in East and possible Central Java. The subduction of the Roo Rise, an area of oceanic basement relief, is thought to contribute significantly to the spatial geochemical source input variations exhibited by Javanese volcanoes.

1. Introduction

Understanding the genesis of volcanic rocks in subduction zone settings is complicated by the multitude of differentiation processes such as fractional crystallisation, magma mixing and

crustal assimilation and geochemical variations in source components, for example, the mantle wedge, subducted oceanic crust and accompanying sediments or melts thereof, that exert control on volcanic rock geochemistry (e.g., Davidson et al., 1987; Hildreth and Moorbath, 1988; McCulloch and Gamble, 1991; Hawkesworth et al., 1991; Pearce and Peate, 1995). Furthermore, within-arc variations in crustal architecture and the nature (age and composition) of the subducting slab and its associated sediments augment the difficulty of disentangling differentiation effects on primary magmas from magma source characteristics. Yet, in order to obtain an accurate understanding of element transfer in subduction zones, it is of primary importance to identify shallow-level geochemical and isotopic modifications of original source compositions in subduction zone petrogenesis.

The Java sector of the Sunda arc in Indonesia is the subject of a long-standing debate on the relative importance of subduction input of crustal material versus magmatic interaction with the surrounding arc crust in controlling the geochemistry of erupted volcanic rocks (e.g., Whitford, 1975; 1982; Wheller et al., 1987; Gasparon and Varne, 1998; Turner and Foden, 2001; Gertisser and Keller, 2003a; Chadwick et al., 2007; Handley et al., 2007; 2011; Halldórsson et al., 2013). Early studies attributed a decrease in Sr isotope composition in lavas from West Java to Bali to diminishing crustal assimilation as a result of eastward arc-crustal thinning (Whitford, 1975; Hamilton, 1979). Gasparon and Varne (1998) also suggested that crustal assimilation accounts for Sr, Pb and Nd isotope systematics along the Sunda arc, from Sumatra to Lombok, relative to mid-ocean ridge basalt (MORB). In addition, more recent detailed geochemical and isotopic studies on individual Javanese volcanoes implicate an important role for crustal assimilation processes in the geochemical and isotopic evolution of magma and its phenocrysts (e.g., Chadwick et al., 2007; Handley et al., 2008a; Abdurrachman, 2012; Abdurrachman and Yamamoto, 2012; Troll et al., 2013). However, addition of a subducted crustal component to the Java sector of the Sunda arc mantle wedge is advocated by others to exert important control on the crustal signature observed within the volcanic rocks (e.g., Whitford, 1982; Wheller et al., 1987; Edwards, 1990; Turner and Foden, 2001; Gertisser and Keller, 2003a; Handley et al., 2007; 2010; 2011). Wheller et al. (1987) suggested that at least three geochemically and isotopically distinct source components are involved in magma genesis: peridotitic mantle, subducted crustal input and a K-group-rich, low-Sr isotope, mantle-derived component. Edwards et al. (1993) used B/Be and radiogenic isotopes of seven volcanic centres along the Sunda and Banda arc (including Guntur and Cereme from the volcanic front in Java) to suggest that the composition of the subduction input is homogenous along the Sunda arc. In contrast, Handley et al. (2011) argued for along-

arc heterogeneity in this component in Java based primarily on Nd-Hf isotope compositions of the volcanic rocks. A recent chemical and isotope (He-C-N) study of active fumarole and hydrothermal gas and water by Halldórsson et al. (2013) proposed that the major volatile budget of the western Sunda arc (Sumatra, Java and Bali) is dominantly sourced from subduction input of crustal material rather than magmatic interaction with thick/old crustal basement.

Considering the nature of the debate, it is surprising that only relatively few Pb isotope data are available for Javanese volcanic rocks. The significant contrast in Pb concentration and isotopic composition of the mantle relative to marine and continental sediments (e.g., Sun, 1980; Price et al., 1986; Ben Othman et al., 1989; Rehkämper & Hofmann, 1997; Gasparon and Varne 1998; Chauvel and Blichert-Toft, 2001) dictates that Pb isotopic ratios of mantle-derived magma are highly sensitive to the addition of subduction-related and/or assimilated arc crustal material during volcanic rock petrogenesis (e.g., James, 1982; Whitford, 1982; Miller et al., 1994; Carpentier et al., 2008). Previously published Pb isotope data of Javanese volcanoes are largely confined to studies analysing only one or two samples from each volcano, with the data amalgamated in broad arc overviews (Whitford, 1982; Turner and Foden, 2001; Woodhead et al., 2001). Prior to this study, only Guntur and Merapi volcanoes have five or more samples with published Pb isotope analyses (Edwards, 1990; Gertisser and Keller, 2003a). Yet, to understand the contribution of crustal assimilation and the nature of assimilants, it is essential to have tight constraints on differentiation characteristics at individual volcanoes.

The oxygen isotope compositions of subduction zone volcanic rocks can also yield key information on crustal assimilation processes during magmatic differentiation and therefore, help to discriminate between subduction-related and assimilated arc crustal inputs. This is due to the significant contrast between the low and relatively constant $\delta^{18}\text{O}$ values in the upper mantle ($+5.5 \pm 0.2\text{‰}$ (Eiler, 2001) to $+5.7 \pm 0.2\text{‰}$ (Harmon and Hoefs, 1995) in MORB relative to Standard Mean Ocean Water (SMOW)) and generally high $\delta^{18}\text{O}$ values of upper crustal materials (typically $>10\text{‰}$; e.g., Ito and Stern, 1986; Davidson and Harmon, 1989) that have undergone low temperature interaction with meteoric water. Oceanic lithosphere may have low and moderately high $\delta^{18}\text{O}$ values (typically between 3-10‰) depending on whether high-temperature hydrothermal alteration or low-temperature alteration and weathering of rocks occurred. Normal depth profiles of oceanic lithosphere show ^{18}O -

enriched pillow lavas and ^{18}O -depleted sheeted dikes and uppermost gabbros (Gao et al., 2006 and references therein).

Here we present new whole-rock Pb isotope data for volcanoes in West Java (Salak, Gede Volcanic Complex and Galunggung), Central Java (Merbabu and Merapi) and East Java (Ijen Volcanic Complex) (Fig.1). New whole-rock oxygen isotope data are presented for Merbabu volcano and new geochemical and radiogenic isotope (Pb, Sr, Nd, Hf) data for three West Javanese Neogene sedimentary rocks of varied lithology are also presented. The new volcano Pb isotope data are combined with previously published geochemical and isotopic data for the same samples (Turner and Foden, 2001; Gertisser and Keller, 2003a; Handley et al., 2007; 2008a; 2010; 2011) and with other published Javanese volcanic rock data to increase our understanding of the relative importance of both crustal assimilation and subduction input of crustal material in Sunda arc volcanic petrogenesis. This study also aims to provide insight into the nature of the underlying arc crust involved in crustal assimilation and thereby expose likely transitions in the composition of the arc basement in Java. We show that through detailed studies of individual volcanoes it is possible to resolve ‘shallow’ versus ‘deep’ crustal inputs in arc magma genesis and we discern the geochemical and isotopic modification of mantle wedge source compositions by both crustal assimilation and subducted crustal input in volcanic petrogenesis in Java.

2. Tectonic and geological setting

The Sunda arc, stretching from the Andaman Islands north-west of Sumatra, through Java to Flores in the Banda sea is part of the Indonesian subduction zone system, formed by subduction of the Indo-Australian plate beneath the Eurasian Plate at a rate of $\sim 6\text{--}7\text{ cm yr}^{-1}$ (Le Pichon, 1968; DeMets et al., 1990; Tregoning et al., 1994; Fig. 1a). The tectonic setting of the Indonesian region is described in detail in Hamilton (1979) and Hall (2002; 2011). The Sunda arc basement is thought to be made up of a series of Gondwanan lithospheric fragments that were accreted to the pre-Cretaceous (Sundaland) Eurasian Plate margin in the Cretaceous (e.g., Metcalfe, 1990; Wakita, 2000; Hall, 2011; Fig. 1a). The associated Cretaceous accretionary-collisional complexes are exposed in Central and West Java (Fig. 1a). The Luk Ulo complex near Karangsambung in Central Java consists of ophiolitic rocks (mafic and ultramafic rocks), sedimentary rocks, and crystalline schists and gneisses occurring as tectonic slabs in a black-shale matrix tectonic *mélange* (Wakita et al., 1994; Kadarusman et al., 2007 and references therein). Such complexes are also exposed at Ciluteh in West Java and Jiwo Hills in Central Java (Metcalfe, 1990; Wakita, 2000; Fig. 1a).

However, the majority of exposed rocks on Java consist of younger, Cenozoic sedimentary sequences and volcanic arc rocks (e.g., van Bemmelen, 1949; Hamilton, 1979; Clements et al., 2009). A major structural feature and geological division is inferred between Central and East Java, running ~NE-SW through the island, close to Merapi at the volcanic front and Muria volcano at the rear arc (Fig. 1b). This lineament is described as either the strike-slip Central Java Fault (Chotin et al., 1984; Hoffmann-Rothe et al., 2001) or the inferred Progo-Muria lineament (Smyth et al., 2005; Smyth et al., 2007) and is suggested to mark the eastern limit of accreted Cretaceous terranes and sutures (Hoffmann-Rothe et al., 2001) or western limit of Archean-aged zircons in the southern mountains (Smyth et al., 2007), respectively. Handley (2006) also proposed a significant crustal boundary at this location (between Merapi and Kelut) based on a compilation of volcanic Sr isotope data, volcano activity and volcano morphology. East of Merapi, the topography along the volcanic front changes from the deeply dissected mountainous and more rugged topography of West and Central Java to the relatively flat East Javan topography, punctuated by large conical volcanoes with relatively smooth volcanic slopes (Handley, 2006; Fig. 1c). Wheller et al. (1987) suggested that the area between Merapi and Kelut is an extinct sector of the arc, containing the volcanoes of Wilis and Lawu, which only display solfataric activity. Recent work highlighting the structural complexity of the Java crust is detailed in Smyth et al. (2007) and Clements et al. (2009).

The subducting plate also shows variability along the arc. The age of the subducting Indian Ocean crust increases from West (~80 Ma) to East Java (~130 Ma) (Fig. 2; Hamilton, 1979; Syracuse and Abers, 2006), while the dip angle of the slab decreases slightly from 50° to 42° (Syracuse and Abers, 2006). The location of the volcanoes in relation to the slab depth, H , and distance from the trench varies quite dramatically along the Java sector of the arc (Figs. 2c and d) but unlike slab age and dip, slab depth does not show a simple increase or decrease along strike (Hutchison, 1982; Syracuse and Abers, 2006). The vertical depth to the slab and the horizontal distance to the trench increase from Krakatau to Ungaran/Dieng volcanoes (~110°E) then abruptly decrease to Wilis, Lawu and Kelut (~112°E) before increasing again to Ijen (114°E). The relatively systematic decrease in slab dip from West to East Java (Fig. 2b) rules out significant lateral slab-tear as an explanation for the variation in slab depth (cf. Garwin et al., 2005). A recent study using Hough Transform analysis of volcano positions in Java suggests that stress regimes in the arc lithosphere control the spatial distribution of the volcanoes (Pacey et al., 2013). Present day collision and subduction of the Roo Rise between 109°E and 115°E (Fig. 1b), an area of oceanic basement relief, has caused dramatic changes in the character of the Java plate margin and a transition from subduction

accretion (west of $\sim 109^\circ\text{E}$ at the Java Trench) to subduction erosion (east of $\sim 109^\circ\text{E}$ at the Java Trench) (Kopp et al., 2006). This is manifest in a marked change from a thick accretionary wedge, continuous outer forearc high and sediment filled trench in the west to no distinct frontal prism or continuous outer forearc high and displacement of the trench and deformation front by ~ 60 km to the north in the east. The northward displacement of the trench is observed between 109°E and 115°E (Kopp et al., 2006), corresponding across strike to volcanoes in Central and East Java. The composition and mass of sediment in the Java Trench also vary along the Java sector. Plank and Langmuir (1998) proposed that 300 m of sediment is subducted beneath Java. Up to 5 km of sedimentary material fills the Sumatra Trench, less than 1 km exists in the western Java Trench and virtually no trench sediments are present in the eastern Java Trench (Plank and Langmuir, 1998; Kopp et al., 2006). The thicker sedimentary deposits present at the site of subduction in West Java, compared with East Java, are a result of the closer proximity of West Java to turbiditic material sourced from the Himalayan collision zone and deep-sea fans surrounding India (Plank and Langmuir, 1998). Sediments deposited on the Indian Ocean Plate south of the trench are relatively uniform in thickness along the arc (200-400 m) (Hamilton, 1979; Moore et al., 1980) and are dominantly detrital-poor, pelagic sediments (Hamilton, 1979).

3. Sample selection and data presentation

To facilitate investigation of differentiation processes and source compositions, samples were selected for Pb isotope analysis from West (Salak and Gede Volcanic Complex), Central (Merbabu) and East (Ijen Volcanic Complex) Javanese volcanoes (Fig. 1c) for which there is no prior published Pb isotope data but for which geochemical and other isotope (Sr-Nd-Hf and some O) data exist (Handley et al. 2007; 2008a; 2010; 2011). To assist in constraining crustal assimilation processes, the sub-sets of samples selected span the full range in SiO_2 content measured at each volcano. New Merapi (Central Java) samples were analysed from the 2006 and 2010 eruptions. Additional samples from Galunggung and Merapi volcanoes were selected that have previously published thermal ionisation mass spectrometry Pb isotope data. These samples were re-analysed in this study to facilitate comparison between data sets collected in different laboratories using different methods and instrumentation (e.g., Tl-doped multi-collector inductively-coupled plasma mass spectrometry (MC-ICP-MS) with sample-standard bracketing data versus un-spiked, conventionally fractionation-corrected thermal ionisation mass spectrometry (TIMS) data). The Pb isotope data comparison is provided in the Appendix along with further details about the selection of previously published Javanese

data sets for Pb isotope data comparison in this study. New whole-rock oxygen isotope data are presented for Merbabu for which there was no previously published O isotope data. The new Javanese volcano Pb and O isotope data are displayed in Tables 1 and 2, respectively. Samples from Gede Volcanic Complex include volcanic rocks from the main Gede edifice, twin volcanic centre Pangrango ($n = 1$), nearby Gegerbentang vent ($n = 1$) and Older Quaternary volcanoes ($n = 1$; Table 1). Samples from Salak include those from both the central vent and flank vents and a Pre-Salak pumice sample (S100). Those from Ijen Volcanic Complex include volcanic rocks from the caldera-rim and intra-caldera eruptive centres (Table 1). Little geochemical and isotopic information is available for upper crustal rocks in West Java. Therefore, three West Javanese upper crustal, Tertiary sedimentary rocks have been analysed (major element and trace element and Pb-Sr-Nd-Hf isotopic compositions; Tables 1 and 3). A calcareous sediment/marl (SED-A), a volcanoclastic sandstone (SED-B) and a mudstone (SED-C) were collected from a river valley near the Bukit Pelangi golf course, approximately 10 km east of Bogor and 20 km NW of the summit of Gede volcano (Fig. 1c, Table 3). The sedimentary units dip between 20-30 degrees to the south-south west, towards Gede volcano. Abundant planktic foraminifera (*Globigerinoides* and *Orbulina*) in the marl suggest a Neogene (likely, upper Miocene or younger) age for the rocks. These data provide information on the likely character of potential crustal assimilants in West Java and particularly at Gede volcano.

The island of Java and the strike of subduction, as indicated by the Java Trench, are oriented approximately east-west. Therefore, the longitude of the volcano is used to represent volcano position along the arc. As across-arc changes in chemistry are recognised at the Sunda arc (Rittman, 1953; Whitford and Nicholls, 1976; Hutchison, 1976; Edwards, 1990) the rear-arc volcanoes of Muria (370 km above the Wadati-Benioff zone (WBZ) in Central Java) and Ringgit Beser (210 km above the WBZ in East Java) (Fig. 1c) are excluded from data comparison. One of the two Merapi samples analysed by Woodhead et al. (2001) displays anomalously low Pb isotope ratios relative to the other Merapi Pb isotope data and to the Central Java group. This sample, therefore, has not been plotted on the figures or considered in the discussion.

To facilitate the identification of geochemical trends, volcanoes are grouped into West, Central and East Javanese provenance (Fig. 1c). Accordingly, the West Java group includes volcanoes from Danau to Cereme, the Central Java group from Slamet to Merapi and the East Java group from Lawu to Ijen (Fig. 1). Slamet, the westernmost Central Java volcano, shows several geochemical similarities to West Java (e.g., lower Ba concentration

and Ba/Hf ratio) yet displays high Sr isotope ratios similar to other Central Java volcanoes and, therefore, has been grouped with Central Java. The placing of Lawu and Wilis is uncertain, whether Central or East Java, as there is limited geochemical and, particularly, isotopic data available. Morphologically, these volcanoes are similar to other East Java volcanoes (single, large composite volcanoes) and lie to the east of the Central Java Fault/Progo-Muria lineament and hence have been placed in the East Java Group. Due to the multiple spellings of Javanese volcano names, those used are consistent with the spellings given the Global Volcanism Program database (<http://www.volcano.si.edu/index.cfm>).

4. Analytical techniques

Lead isotope ratios for Salak, Gede, Galunggung, Merapi and Ijen volcanic rocks and the three West Java sedimentary rocks were determined on bulk-rock powders by wet chemistry and multi-collector inductively-coupled plasma mass spectrometry (MC-ICP-MS; Nu Plasma 500 HR) using Tl doping and sample-standard bracketing (White et al., 2000) at the Ecole Normale Supérieure in Lyon. Details of the preparation of rock powders can be found in Turner and Foden (2001), Gertisser and Keller (2003a), Handley et al. (2007; 2008a; 2010; 2011) and Preece et al. (2013). The whole-rock powders were first leached in hot 6 M HCl prior to attack in a 3:1:0.5 mixture of concentrated HF:HNO₃:HClO₄. The samples were then taken up in 6 M HCl after fuming with HClO₄ to eliminate fluorides. Lead was separated by ion-exchange chromatography on 0.5 mL columns filled with Bio-Rad AG1-X8 (100-200 mesh) resin using 1 M HBr to first elute the sample matrix followed by 6 M HCl to subsequently recover the Pb. The total procedural Pb blank was < 20 pg. The NIST 981 Pb standard and the values of Eisele et al. (2003) were used for bracketing the unknowns (every two samples), and Tl was used to monitor and correct for instrumental mass bias. Internal uncertainties on the reported Pb isotope ratios are 50-100 ppm. The Pb isotope data are listed in Table 1.

The Merbabu Pb isotope measurements were carried out on bulk-volcanic rock powders without further pre-treatment at the University of Tübingen using analytical methods described in detail by Hegner et al. (1995a, 1995b). The preparation of rock powders follows that detailed in Gertisser and Keller (2003a). Lead was separated on Teflon columns containing 80 µl AG 1-X8, 100-200 mesh and employing a HBr-HCl wash and elution procedure. Lead isotope compositions were determined by thermal ionisation mass spectrometry on a Finnigan MAT 262 mass spectrometer in static collection mode. Lead was loaded with a Si-gel onto a pre-conditioned Re filament and measured at 1250-1300°C. Total

procedure blanks were < 50 pg. A factor of 0.1% fractionation per mass unit was applied to all Pb isotope analyses, using NBS SRM 982 as the reference material and NBS SRM 981 as a standard. The estimated 2 σ uncertainty of the reported Merbabu volcano Pb isotope ratio is better than 0.01 %.

Merbabu whole-rock oxygen isotope ratios were measured using a Finnigan MAT 252 gas source mass spectrometer (Hegner et al., 1995b). Silicate oxygen was extracted by bromine pentafluoride at 600°C, followed by conversion to CO₂. The results are reported as per mil (‰) deviations relative to Standard Mean Ocean Water (SMOW) and refer to a certified value of 9.6 \pm 0.1‰ measured on NBS 28. For the O isotope determinations at least two measurements were performed for a given sample and the average $\delta^{18}\text{O}$ values are reported in Table 2. All values were reproduced within an analytical error of 0.1-0.2‰.

The major element, trace element and Sr–Nd–Hf isotopic data for the West Java sedimentary rocks were collected using the same procedures and data quality constraints as those given in Handley et al. (2010) and Handley et al. (2011). Full details on the analytical techniques for the West Java sedimentary rocks are provided in the Appendix.

5. Results

5.1. Pb isotope data

The new Pb isotope data for volcanic rocks from Salak, Gede Volcanic Complex, Galunggung, Merbabu, Merapi and Ijen Volcanic Complex are presented in Table 1 and Fig. 3. The new data (colour-filled symbols in Figs. 3a and b) form a positive array in $^{207}\text{Pb}/^{204}\text{Pb}$ and $^{208}\text{Pb}/^{204}\text{Pb}$ versus $^{206}\text{Pb}/^{204}\text{Pb}$ space and display higher Pb isotope ratios than Indian Ocean mid-ocean ridge basalt (I-MORB) and seamounts subducting at the Java Trench, represented by the Eastern Wharton Basin and Argo Basin Volcanic Provinces (I-Seamount East). Most of the Javanese volcanic rock data overlap with the fields for local Indian Ocean subducted sediments (I-SED), Indian Ocean Mn crusts and to a lesser extent, the limited data available for the local upper crust (Sumatran intrusive rocks (Intrusive), calcareous sediment upper crust samples from Central Java and the new West Java upper crust samples (Local Crust)). Ijen volcanic rocks have the most primitive Pb isotopic compositions, with most displaying less radiogenic ratios than local subducted sediment and local upper crustal sedimentary rocks (Local Crust). The Gede volcanic rocks exhibit the most radiogenic values for Javanese volcanoes so far determined (Figs. 3c and d), with the associated centres of Pangrango and Gegerbentang displaying the highest Pb isotopic ratios (Table 1). Salak and Galunggung show a similar range in Pb isotope ratios, overlapping with the more primitive

end of the Gede data array. The new data for Merapi ($n = 13$, Table 1) which consists of both high-K (< 1900 ^{14}C y B.P.) and medium-K (> 1900 ^{14}C y B.P.) rock types (Gertisser and Keller, 2003b) and recently erupted 2006 and 2010 volcanic rocks, show remarkable within-suite homogeneity in Pb isotopic composition compared to the other volcanoes. However, the Merapi rocks lie slightly elevated, relative to the main Java trend, at higher $^{207}\text{Pb}/^{204}\text{Pb}$ for a given $^{206}\text{Pb}/^{204}\text{Pb}$ (Fig. 3c). The Pb isotope ratios of Merbabu volcanic rocks from Central Java overlap with the data for Salak and Gede.

The local, upper crustal, sedimentary rocks from West Java display relatively similar Pb isotopic compositions to one another and lie at intermediate Pb isotopic ratios relative to the Javanese volcanic rocks (Figs. 3c and d). The sedimentary rocks exhibit more primitive isotopic ratios than Gede Volcanic Complex rocks and more radiogenic values than Ijen Volcanic Complex. In $^{208}\text{Pb}/^{204}\text{Pb}$ - $^{206}\text{Pb}/^{204}\text{Pb}$ space, the marl and mudstone rocks lie slightly off the main Javanese volcanic rock trend, at lower $^{208}\text{Pb}/^{204}\text{Pb}$ for a given $^{206}\text{Pb}/^{204}\text{Pb}$.

5.2. Oxygen isotope data

New bulk-rock oxygen isotope data of Merbabu volcano are presented in Table 2 and Fig. 4. Merbabu $\delta^{18}\text{O}$ values range from +6.4 to +8.4‰, are higher than typical values for MORB of $+5.5 \pm 0.2$ ‰ (Eiler, 2001) and $+5.7 \pm 0.2$ ‰ (Harmon and Hoefs, 1995) and extend to the highest volcanic whole-rock $\delta^{18}\text{O}$ values yet reported for Java. The Merbabu $\delta^{18}\text{O}$ values largely overlap with the range in whole-rock values reported for neighbouring Merapi volcano and Cereme, West Java (insets to Figs. 4a and b), which also display higher overall $\delta^{18}\text{O}$ values relative to $\delta^{18}\text{O}$ bulk volcanic rock data of Galunggung (Figs. 4a and b). Low loss on ignition (LOI) values in the Merbabu volcanic rocks of -0.44 to 0.71 wt%, except for MB-6 with a higher LOI value of 1.12 wt% (Handley et al., 2011), suggest that post-eruption secondary alteration processes, such as weathering, have not significantly affected the rocks and by inference, the measured oxygen isotope values. The samples with the highest $\delta^{18}\text{O}$ values (MB-16 and MB22) do not correspond to the sample with the highest LOI (MB-6). The Merbabu $\delta^{18}\text{O}$ values are much lower than those of the local calcareous sedimentary rocks given by Gertisser and Keller (2003a) (+18.9 to 20.5‰), and local limestone samples of Troll et al. (2013) (+24.0 to 24.5‰), and also lower than values measured in local volcanoclastic crust (+12.5; Troll et al., 2013) and bulk calc-silicate xenoliths found within Merapi lavas (+11.5 to 14.2‰; Gertisser and Keller, 2003a; Troll et al., 2013). Note that the calc-silicate xenolith $\delta^{18}\text{O}$ values likely represent the resultant limestone-magma interaction

rather than the carbonate protolith. The highest Merbabu $\delta^{18}\text{O}$ values correspond to the samples with the lowest Pb isotope ratios (Fig. 4a inset).

6. Discussion

It is widely considered that the majority of Javanese arc magmas are formed through flux-melting of the mantle wedge due to subduction of the Indo-Australian Plate beneath the Eurasian plate (e.g., Harmon and Gerbe, 1992; Gertisser and Keller, 2003a; Handley et al., 2007; 2011). Although, the low volatile contents measured in mafic glass inclusions in high-magnesium basalts from Galunggung, led Sisson and Bronto (1998) to suggest that pressure-release melting of hot mantle peridotite may play a role in magma genesis for some Javanese volcanoes. Upon ascent to the surface, the major and trace element composition of magmas may be modified by a number of processes such as fractional crystallisation, magma mixing and crustal assimilation (e.g., Anderson, 1976; Sakuyama, 1979; DePaolo, 1981; Thirlwall et al., 1996). Yet, the radiogenic isotopic composition of a magma is not expected to change by recent partial melting and fractional crystallisation or magma mixing, provided that the mixed magmas are from the same source. However, the isotopic ratios of magma are susceptible to modification by assimilation of crustal material during ascent and shallow-level storage, if the magma and assimilated material are characterised by contrasting isotopic ratios (e.g., Davidson et al., 2005). Similarly, stable O isotope values of mantle-derived melts may be significantly modified by assimilation of crustal material that has experienced low-temperature interaction with meteoric water or seawater, or interaction with high-temperature hydrothermal fluids (e.g., James, 1981; Bindeman et al., 2001). Therefore, the new Pb and O isotope data presented here can be used to elucidate the relative role that crustal assimilation has exerted on magma chemistry during differentiation of Javanese rocks as a means to “see through” to source input variations.

6.1. Pb isotopic evidence for assimilation of crustal material

Owing to the large contrast in Pb concentration and Pb isotopic composition between mantle-derived magma (low concentration and less radiogenic) and continental upper crust (high concentration and more radiogenic) the former is sensitive to incorporation of crustal material during differentiation in the arc crust (e.g., Davidson, 1987). If assimilation is concomitant with fractional crystallisation of magma (AFC) (e.g., DePaolo, 1981), evidence for assimilation may be visible through correlations of isotope ratios with indices of differentiation (e.g., SiO_2 , MgO).

The plot of $^{207}\text{Pb}/^{204}\text{Pb}$ versus SiO_2 content for Javanese volcanic rocks (Fig. 5) highlights the significantly higher $^{207}\text{Pb}/^{204}\text{Pb}$ for West (Gede, Salak, Galunggung) and Central (Merbabu, Merapi) Java compared to East Java (Ijen) at similar degrees of magmatic differentiation. The Pb isotope ratios of Ijen volcanic rocks do not correlate with indices of differentiation, though they do show some degree of scatter. The volcanic rocks from Salak show no clear correlation despite inferred crustal assimilation at this centre based on the modelling of trace elements and Sr isotope ratios of the Central Vent Group (CVG) (Handley et al., 2008a). This may be due to the slightly more limited number of CVG samples analysed for Pb isotopes ($n = 3$; Table 1) compared to those analysed for Sr isotopes ($n = 5$). Negative correlations between Pb isotope ratios and SiO_2 are observed for Merbabu ($R^2 = 0.85$) and Gede Volcanic Complex ($R^2 = 0.71$), suggesting the incorporation of isotopically contrasting material at these volcanoes during magmatic differentiation. The correlations observed for Merbabu and Gede show that the lowest SiO_2 content rocks, inferred to represent the least differentiated samples, have the most radiogenic Pb isotopic compositions. The three analyses available for Galunggung also suggest a similar, negative trend. Two possible hypotheses for the observed negative trends are: 1) increasing incorporation of less radiogenic material, such as more isotopically primitive arc rocks or ophiolitic oceanic crust (now in the arc crust), during progressive magmatic evolution or, 2) the lowest silica rocks have assimilated the most (by volume) radiogenic crustal material. The exposed arc basement in West and Central Java is composed of accreted Mesozoic ophiolitic and arc rocks and Eocene to Late Miocene volcanogenic turbidites and breccias, quartz-rich sandstones and limestone (Clements and Hall, 2007; Section 2). The observed negative Pb isotope trends of Gede and Merbabu volcanic rocks may, therefore, be explained by hypothesis one, whereby progressive assimilation of less radiogenic mafic-ultramafic (ophiolitic-type) or more primitive arc basement occurs during magmatic differentiation (Fig. 5). A similar model has been proposed for Lopevi volcano in the intra-oceanic Vanuatu arc, where negative correlations between Sr isotope ratios and SiO_2 content are observed (Handley et al., 2008b). With regard to the second hypothesis, there are two further possibilities: a) the higher temperature and lower viscosity of the more primitive, lower SiO_2 magma enabled assimilation of a greater volume of crust, relative to more evolved magma (e.g., Huppert and Sparks, 1985; Peccerillo et al., 2004) or b) the negative Pb isotope correlation observed for Gede and Merbabu is explained by assimilation of a component characterised by low SiO_2 content and high Pb isotope ratios such as limestone/calcareous sediment. Assimilation of limestone has already been suggested as an important process during magmatic evolution at Merapi volcano (Chadwick et al., 2007;

Deegan et al. 2010; Troll et al., 2013), adjacent to Merbabu. Limestone sampled from a carbonate platform at Parangtritis, south of Merapi, has low SiO₂ (0.28 wt %), Na₂O (0.12 wt %) and K₂O (0.00 wt %) contents and high CaO and CO₂ contents (Deegan et al., 2010) but unfortunately, Pb isotope data are not available for these samples. However, the Pb isotope composition of a local, low SiO₂ content, West Javanese calcareous marl (SED-A, Tables 1 and 3) collected 20 km NE of the summit of Gede volcano in West Java and two calcareous marls collected from the Wonosari Beds, Southern Mountains, south-east of Imogiri (Miocene) and Djiwo Hills, south of Klaten (Eocene), relatively close to Merbabu and Merapi in Central Java (Gertisser and Keller, 2003a) have lower Pb isotope ratios than the majority of West and Central volcanic rocks (Figs. 3 and 5). Therefore, crustal assimilation of the local calcareous sedimentary rocks (analysed to date) cannot explain the high Pb isotope ratios of the West and Central Javanese rocks in the lowest SiO₂ samples (Fig. 5) or the observed correlation between ²⁰⁷Pb/²⁰⁴Pb and SiO₂.

To further investigate the potential control of carbonate assimilation on the whole-rock geochemical composition of Javanese volcanic rocks, experimental research carried out on carbonate assimilation at Italian volcanoes can be utilised. Experimental work on the interaction between limestone and magma in connection with Italian volcanoes has shown that assimilation of limestone creates a negative correlation between Na₂O + K₂O and SiO₂, or a horizontal array at low SiO₂ contents. The experimental data trends (shown as arrows on Fig. 6) agree well with melt inclusion data of volcanic rocks at Vesuvius volcano (Iacono Marziano, 2008). Figure 6 shows an overall positive correlation in Na₂O + K₂O versus SiO₂ space for the Javanese data set and, more importantly, a strong positive correlation for Gede ($R^2 = 0.76$) and a slightly weaker correlation for Merbabu ($R^2 = 0.58$). This suggests that, unlike Vesuvius, limestone assimilation or assimilation of local calcareous sediments (SED-A, Table 3) does not fully control the major element composition of most Javanese volcanic rocks. Furthermore, the range in silica content observed at Gede and Merbabu is comparable to the other Javanese volcanoes (e.g., Ijen) and is not significantly lower as might be expected if significant limestone/carbonaceous sediments were being assimilated. The Central Javanese volcanoes do show relatively more scatter in intra-volcanic trends and higher Na₂O + K₂O for a given SiO₂ content compared to West Javanese rocks suggesting a potential possible role for some degree of carbonate assimilation in Central Java relative to West Java but does not dominate the whole-rock geochemical composition of the erupted rocks. Alternatively, the Central Javanese volcanoes are located at greater distance from the trench and subducting slab (Fig. 2), which may explain their generally higher K₂O content.

The new Pb isotope data of Merapi volcanic rocks, provide two surprising observations. Firstly, that there is little difference in the Pb isotope ratio between the temporally divided medium-K and high-K rock groups (Table 1; Gertisser and Keller, 2003a). Secondly, that despite strong evidence for crustal contamination by carbonate/more radiogenic material from the Sr isotope, crystal stratigraphy study of Chadwick et al. (2007), the volcano displays an extremely limited within-suite range in whole-rock Pb isotopic composition. This is particularly noticeable when compared to the within-suite range observed at Gede, Salak and Ijen. The Merapi samples lie at higher Pb isotope compositions than the available Pb isotope data for local calcareous crustal rocks (Figs. 3 and 5) and lie slightly off the main Java array, at elevated $^{207}\text{Pb}/^{204}\text{Pb}$ for a given $^{206}\text{Pb}/^{204}\text{Pb}$. Greater knowledge of the Pb isotopic compositional range of the local crust, including carbonaceous sediments is required in order to fully determine the cause of the displacement of Merapi data from the main Java trend an apparent buffering of the volcano's Pb isotope composition.

In summary, the Pb isotope data show evidence for crustal assimilation processes at Gede and Merbabu (and possibly Galunggung). The negative correlation between Pb isotope ratio and SiO_2 content suggests either that the more mafic, higher temperature rocks have assimilated the most (by volume) radiogenic crustal material, or that the crustal assimilant is characterised by a lower Pb isotopic composition relative to the source magma. These options are discussed further below using constraints from O isotopes and trace element data. The Pb isotope compositions of the volcanic rocks and local crustal rocks from West and Central Java, combined with $\text{K}_2\text{O} + \text{Na}_2\text{O}$ versus SiO_2 volcanic data trends suggest little importance of the assimilation of carbonaceous material at Gede. The overall higher $\text{K}_2\text{O} + \text{Na}_2\text{O}$ contents and more scattered data trend in $\text{K}_2\text{O} + \text{Na}_2\text{O}$ - SiO_2 space for Merbabu, relative to Gede, suggest potentially a limited role for the assimilation of carbonaceous material. However, to date, the Pb isotope ratios available for local carbonaceous crustal rocks do not provide high enough Pb isotope ratios to be suitable contaminant end members to support this model. The lower Pb isotopic ratios in Ijen volcanic rocks, compared to Central and West Java, do not correlate with indices of differentiation indicating that assimilation of isotopically distinct crustal material is not an important process in modifying magma isotopic compositions at this volcano.

6.2. Oxygen isotope constraints on crustal assimilation

Oxygen isotope data can be utilised to further investigate the suggested importance of crustal assimilation at Merbabu and Gede gained from Pb isotope data. The new Merbabu whole-

rock $\delta^{18}\text{O}$ data are significantly higher than typical values for MORB ($+5.7 \pm 0.2\text{‰}$, Harmon and Hoefs, 1995 and $+5.5 \pm 0.2\text{‰}$, Eiler, 2001) (Table 2; Fig. 4) suggesting possible interaction between magma and upper crustal material that has undergone low-temperature interaction with meteoric water. The Merbabu $\delta^{18}\text{O}$ values are similar to the elevated whole-rock $\delta^{18}\text{O}$ values of Merapi, which also displays elevated plagioclase and clinopyroxene $\delta^{18}\text{O}$ values (Fig. 4) and for which crustal assimilation processes have been implicated (Gertisser and Keller, 2003a; Chadwick et al., 2007). The two highest $\delta^{18}\text{O}$ values are found in the lowest Pb isotope rocks (Fig. 4a) suggesting that the assimilant may be characterised by lower Pb isotope ratios than the ascending magma.

The previously published olivine and clinopyroxene O isotope mineral separate data from Gede Volcanic Complex possess relatively restricted $\delta^{18}\text{O}$ values and lie largely within error of mantle $\delta^{18}\text{O}$ values (Figs. 4b and c; Handley et al., 2010). These values suggest that the assimilant involved has not significantly interacted with low-temperature fluids and, therefore, assimilation may occur relatively deep in the crust, and/or, that the assimilant is characterised by low $\delta^{18}\text{O}$ values. The latter is consistent with assimilation of more primitive arc rocks and/or ophiolitic material, which was suggested as a possible explanation for the negative correlation between Pb isotope ratios and SiO_2 contents of the volcanic rocks. The Gede samples for which both clinopyroxene O isotope data and whole-rock Pb data are available show no correlation between stable and radiogenic isotopes (Figs. 4a and 4c).

Although mineral O isotope data are not available for Merbabu lavas, taking all published Javanese plagioclase and clinopyroxene $\delta^{18}\text{O}$ values into account, Figure 4b shows that there is a significant along-arc variation in $\delta^{18}\text{O}$ for each mineral type, with the highest plagioclase and clinopyroxene $\delta^{18}\text{O}$ values in rocks from Central Java relative to West and East Java. The low values in East Java and lack of correlation between Pb isotope ratios and indices of differentiation at Ijen further suggest minimal modification of magma source compositions via interaction with crustal material during shallow level magmatic differentiation. However, for West Java, the low $\delta^{18}\text{O}$ values may be deceptive when trying to identify crustal assimilation if a low- $\delta^{18}\text{O}$ assimilant is involved. Interaction with hot ^{18}O -depleted meteoric groundwater in hydrothermal circulation at the margins of the magma reservoir has been suggested from $\delta^{18}\text{O}$ values and the geochemical characteristics of lavas at Galunggung in West Java (Harmon and Gerbe, 1992). Salak and Guntur display $\delta^{18}\text{O}$ clinopyroxene values lower than the mantle range (Fig. 4c) likewise suggesting potential involvement of an ^{18}O -depleted component. For Merapi, the wide range in clinopyroxene

$\delta^{18}\text{O}$ values (5.1 to 7.2‰), both above and below the mantle range, may indicate that both ^{18}O -enriched and ^{18}O -depleted components are involved in its petrogenesis. New laser fluorination O isotope mineral data for Guntur volcano also implicate the involvement of ^{18}O -enriched and ^{18}O -depleted components in volcanic petrogenesis (Macpherson et al., submitted). The along-arc peak in $\delta^{18}\text{O}$ values at Central Java contrasts with the along-arc peak in Pb isotope ratios, observed for West Java (Fig. 7a). However, the along-arc trend in oxygen isotopes is similar to the along-arc pattern in Sr isotope composition for which there is significantly more data available (Fig. 7b). This suggests that the main crustal assimilant involved in West Java volcanic petrogenesis is characterised by relatively low $\delta^{18}\text{O}$ and low Pb and Sr isotopic ratios. Whereas in Central Java this component may also be present but an additional assimilant characterised by higher $\delta^{18}\text{O}$ values, higher Sr isotope ratio but relatively conservative Pb isotope ratios, exerts significant control on the isotopic composition of the erupted rocks.

The combined use of O and Pb isotopes can help to constrain the relative importance of crustal assimilation versus subduction input of crustal material in magma genesis and evolution due to the contrasting mixing trajectories that result from each process (e.g., James, 1981; Davidson et al., 2005). The inset diagram of Fig. 4c shows the different curvatures expected for crustal assimilation (dashed lines) and subduction input of crustal material to a mantle wedge source (solid lines) that arise due to the large difference in the Pb/O ratio of the mantle wedge source and the mantle-derived arc magma. Mixing between the mantle source and subducted crustal material results in strongly convex-downward mixing curves (Fig. 4c model A and inset diagram). Whereas mixing between mantle-derived arc magma and the arc crust creates largely straight or convex-upwards curves (James, 1981) (Fig. 4c model B and inset diagram). Acknowledging that the exact locations of the mixing trends in Pb-O isotope space are somewhat dependant on the Pb isotope ratio of the selected crustal end-member (Fig. 4c inset), the Gede and other available Javanese clinopyroxene $\delta^{18}\text{O}$ data generally lie along the mixing curve representing input of local subducted sediment to the mantle source (Fig. 4c model A), opposed to the mixing of arc magma with local arc crust (Fig. 4c model B; see figure caption for the end member composition details). The observed scatter in Gede clinopyroxene $\delta^{18}\text{O}$ values, some of which are higher than the mantle $\delta^{18}\text{O}$ range, may indicate the extent of modification of initial arc magma clinopyroxene $\delta^{18}\text{O}$ values by crustal assimilation processes. However, this could also represent slight variations in the Pb isotopic composition and/or concentration of the subducted crustal component and therefore, the

location of the bend in the calculated mixing curve. For Guntur and Salak, clinopyroxene $\delta^{18}\text{O}$ values lower than the mantle range and below the mixing curve support some involvement of an ^{18}O -depleted component in magma genesis as discussed above. Nevertheless, the modelling suggests that the dominant control on O-Pb isotope systematics at Gede and other West Javanese volcanoes is related to the source input of crustal material.

6.3. Pb isotope constraints on the source components involved in magma genesis

Despite evidence for some modification of initial source Pb isotope ratios via crustal assimilation, the wide contrast in Pb isotope ratios of East Javanese volcanic rocks (e.g., Ijen) compared to those from West and Central Java (e.g., Gede, Salak and Merbabu) at similar degrees of differentiation (Fig. 4) points towards heterogeneity in the source isotope composition, whether in the mantle wedge source component itself or variation in the type and/or amount of the subducted input. Having established that the West Javanese volcanic rocks exhibit higher Pb isotope ratios than the local crustal rocks and that the highest Pb isotope ratios are observed in the least differentiated (inferred, least crustally contaminated) rocks (Fig. 4), the higher Pb isotope ratios of West and Central Javanese volcanic rocks, relative to East Javanese rocks, may represent a greater proportion of subducted crustal material or a similar proportion of more radiogenic material in West and Central Javanese rocks. Handley et al. (2011) showed that Hf-Nd isotope systematics in Javanese lavas are consistent with heterogeneity in the subduction component input along the arc, largely controlled by observed present-day spatial variations in the sediments deposited in the Java Trench. The authors implicate the incorporation of a dominantly continental-derived, detrital-rich subducted sedimentary component in the west and a more pelagic, clay-rich subducted sedimentary component and possibly stronger slab-fluid imprint in the east. The significantly higher radiogenic Pb isotope ratios measured in West Java volcanic rocks, relative to East Java at a similar degree of magmatic differentiation (regardless of whether the highest or lowest Pb isotope ratios of Gede, for example, are taken), are also consistent with a higher proportion of old, detrital-rich continental material in the source in West Java. This idea is explored further in Section 6.4.

The relatively tight Java array displayed in Pb-Pb isotope space can be utilised to provide constraints on the mantle wedge source composition involved in magma genesis. The Java data array projects back towards a source composition with approximately average I-MORB composition ($n = 43$, Fig. 3a and b) with little or no requirement for an Indian Ocean seamount-source component (I-Seamount East; Fig. 3b). The tight, linear array suggests a

relatively homogenous mantle source composition for the Javanese volcanoes presented. This contrasts with the suggestion by Handley et al. (2007) that there may be potential heterogeneity in this component. Handley et al. (2007) implicated a mantle source composition with lower than average $^{176}\text{Hf}/^{177}\text{Hf}$ for Ijen (and by inference East Java) relative to Central and West Java. However, an average I-MORB source Pb isotopic composition is sufficient to explain the overall Java Pb isotope data.

On a Pb-Pb isotope diagram, simple mixing between two components will produce a straight line. The new and previously published data for Javanese volcanoes (excluding Merapi volcano) appear to lie on a simple binary mixing line between average I-MORB, representing the mantle source, and bulk Indian Ocean sediment (I-SED) exemplified by the mixing line between I-MORB and the bulk Java subducted sediment composition of Plank and Langmuir (1998) shown in Figures 3a and b. However, several of the local upper crust rocks, represented by Sumatran intrusive rocks and sedimentary rocks from Central Java, overlap with the field for I-SED. Therefore, from the Pb-Pb diagram alone, the exact nature of the crustal component involved during petrogenesis is not discernable.

It is more difficult to determine the role of a slab fluid produced by dehydration of the down-going altered oceanic crust (AOC) from Pb isotopes. Hydrothermal alteration processes at the ridge-crest may result in a significant amount of Pb being extracted from the oceanic crust prior to subduction (e.g., Albarède and Michard, 1989) minimising the imprint of any addition of this component to the mantle source. However, an estimate of the Pb isotopic composition of an AOC liberated fluid can be gained by using the compositions of Indian Ocean ferromanganese crusts (Mn nodules) (O’Nions et al., 1998; Frank and O’Nions, 1998). Manganese nodules scavenge and incorporate trace metals from seawater and hence record the isotopic composition of ambient seawater (e.g., Goldstein and O’Nions, 1981; Albarède and Goldstein, 1992; Frank and O’Nions, 1998) that would have interacted with I-MORB. Therefore, the Pb isotopic composition of the AOC fluid component is likely to lie on a mixing line between I-MORB and the composition of Mn nodules in Pb-Pb isotope space. As can be seen from Figures 3a and b, the field for 0-20 Ma old Indian Ocean Mn nodules overlaps almost completely with the I-SED field and several upper crust sedimentary rocks and, therefore, this component cannot from its Pb isotope composition alone be distinguished from subducted sediment input or crustal assimilation. The involvement of an AOC component will be discussed further in Section 6.4 using a combined Pb isotope-trace element ratio approach. The Pb isotope composition of the altered oceanic crust end-member used in modelling Sunda and Banda arc volcanic petrogenesis by Edwards et al. (1993) lies at a

reasonable position in $^{208}\text{Pb}/^{204}\text{Pb}$ - $^{206}\text{Pb}/^{204}\text{Pb}$ space (Fig. 3b). However, the proposed $^{207}\text{Pb}/^{204}\text{Pb}$ composition of this component appears to be significantly low, lying on the northern hemisphere reference line (NHRL; Hart, 1984) and not on a mixing line between I-MORB and the Mn crusts. This is probably due to the fact that Edwards et al. (1993) used a generic estimated AOC composition and did not use local end-member (Indian Ocean) compositions, presumably due to the lack of published altered Indian MORB data. We suggest that their $^{207}\text{Pb}/^{204}\text{Pb}$ ratio is unlikely to be representative of the true $^{207}\text{Pb}/^{204}\text{Pb}$ ratio of AOC fluid at the Sunda arc.

6.4. Combined Pb isotopic and trace element constraints on subduction input variability

Barium concentrations in Javanese volcanic rocks show considerable variation along the island, with East and Central Java (excluding Slamet) displaying significantly higher concentrations than those in West Java for a given SiO_2 content (Fig. 8a). Despite some scatter in individual volcanic suites for Central and East Java, the differentiation trends for the geographic suites form largely parallel, positive arrays (noting the exceptionally high Ba concentrations of some Lamongan and Tengger Caldera volcanic rocks) that do not converge upon a common parental composition when projected backwards to less evolved SiO_2 contents. This suggests that the differences in Ba concentrations of the Javanese rocks result from heterogeneity in the source region rather than from shallow-level assimilation processes during magmatic evolution.

Barium, as a large ion lithophile element (LILE), is considered highly mobile in subduction-related fluids relative to the more fluid-immobile, light rare earth elements (LREE) and the high field strength element (HFSE) Hf (Hawkesworth et al, 1993; Keppler, 1996; Kogiso et al., 1997; Kessel et al., 2005). Therefore, Ba/Hf ratios combined with the Pb isotope compositions of the Javanese volcanic rocks may yield potential insight into variations in subduction-related components along the island. Higher Ba/HFSE (e.g., Ba/Hf, Ba/Nb) and Ba/LREE (e.g., Ba/La) ratios in arc lavas, relative to depleted upper mantle values, are usually attributed to significant input of an altered oceanic crust fluid component (e.g., Elliott et al., 1997; Woodhead et al., 2001) but equally may result from the contribution of hydrous melts of subducted sediment (e.g., Kelemen et al., 2003; Hermann and Rubatto, 2009; Jicha et al., 2010). High Ba concentrations, up to 3500 ppm, are reported in drilled Indian Ocean sediments (Gasparon and Varne, 1998). Figure 8b highlights the clear geographic differences of volcanic rocks in West (~high Pb isotope ratios and relatively low Ba/Hf), Central (moderately high Pb isotope ratios and high Ba/Hf) and East (relatively low

Pb isotope ratios and moderate to high Ba/Hf) Java. The boxed arrows on Figure 8b show the estimated maximum modification of Gede source Ba/Hf and Pb isotope ratios due to crustal assimilation. We anticipate a relatively similar, or smaller impact from crustal assimilation on the source Ba/Hf ratios and Pb isotopes of the Central Javanese rocks (Merapi and Merbabu) due to 1) their more restricted range in Pb isotope ratio, 2) the low Ba/Hf ratios and low Pb isotopic ratios of local carbonaceous sedimentary rocks (e.g., Table 3; Fig. 5) and 3) that Slamet volcano, which also displays similarly high Sr isotope ratios compared to the other Central Javanese volcanoes, has low Ba/Hf ratios similar to the West Javanese volcanic rocks (Fig. 7). Nonetheless, regardless of whether the highest or lowest Pb isotope ratio for Gede or Merapi/Merbabu is taken as the rock least affected by crustal assimilation, the geochemical contrasts between the geographic regions still remain.

Simple bulk-mixing models between an I-MORB source and Indian Ocean sediment end-members (see Fig. 8 caption for details) show that the differences in isotopic and trace element ratios between West and East Java can be explained via heterogeneity in the subducted sediment contribution, largely reflecting present-day spatial variations in sediment compositions on the down-going plate in the Java Trench. A low or high Ba/Hf and higher Pb isotopic composition (e.g., detrital-rich) sediment is required for West Java versus a high Ba/Hf, low Pb isotopic composition (e.g., pelagic, clay-rich) sediment is required for East Java. This suggestion of heterogeneity in the subducted sediment component along Java is similar to that proposed by Handley et al. (2011) using Nd-Hf isotopic compositions of Javanese volcanic rocks. If sediment melts/fluids are involved, rather than bulk-sediment addition, the subducted sediment percentage contributions required are smaller than those shown in Figure 8.

An alternative hypothesis to the involvement of a heterogeneous subducted sedimentary component is assimilation of a heterogeneous sedimentary component in the crust. Clements et al. (2009) suggested that the Late Eocene to Early Miocene basin-filling sediment composition varies along Java: West and Central Java basins were supplied with quartz-rich clastic sediments by rivers draining the Sunda Shelf whereas East Java was largely supplied by Eocene to Miocene volcanoclastic products of the Southern Mountains Arc. Crustal assimilation of old, continentally-derived sediments, presumably with radiogenic Pb isotope compositions in West Java, relative to East Java, may explain the eastward decrease in Pb isotope ratios along the island. However, the low, mantle-like $\delta^{18}\text{O}$ values of mineral separates from West Java volcanic rocks would require relatively deep assimilation of the basin-fill sediments, prior to significant magmatic differentiation to avoid 1) O-isotope

modification of the assimilant by low-temperature meteoric fluid, and 2) the detection of assimilation of this component utilising radiogenic isotope and SiO₂ variations. Therefore, we prefer a model of subducted sediment involvement to explain the contrasts in Pb isotope ratios along Java. Furthermore, the addition of arc crust to a magma has less leverage on the Pb concentration (and thereby Pb isotope ratio) than addition of subducted crustal components to the mantle wedge source, due to the significantly low concentration of Pb of the mantle wedge. Therefore, the impact of crustal assimilation will likely be less than the impact of source input of crustal material upon the resultant Pb isotope ratios. This conclusion is consistent with a recent study on the chemical and isotopic (He-C-N) characterisation of active fumaroles and hydrothermal gases and waters from the summits and flanks of multiple volcanic centres along the western Sunda arc (Halldórsson et al., 2013), which suggested that subduction-related source contamination plays the dominant role over thick/old crustal basement in supplying the major volatile output budget of the western Sunda arc volcanoes.

The higher Ba/Hf ratios in East Javanese and possibly also Central Javanese rocks may be attributed to either greater contribution of high-Ba/Hf sediment (fluid/melt) and/or the more significant involvement of an AOC fluid component compared to West Java. A melt inclusion study of Ijen and Tambora volcanic rocks (Vigouroux et al., 2012) suggested that Ijen records a higher AOC-derived fluid component relative to Galunggung and Tambora, with AOC being the main source of Sr and volatiles and subducted sediment melt being an important component for Ba, Pb, Th and the LREE. Although we note that Vigouroux et al. (2012) did not take into consideration the impact of crustal assimilation in their study. An AOC and/or Roo Rise (I-Seamount East) lithospheric liberated fluid component is likely to have less radiogenic Pb isotope compositions than the Javanese volcanic rocks (Section 6.3, Fig. 3) and, therefore, mixing alone between I-MORB and AOC or subducted lithospheric fluid is not sufficient to explain the volcanic data. However, we propose that subduction of the Roo Rise south of Central and East Java (~109°E to 115°E at the trench, ~110°E to 116°E beneath Java; Fig. 1) has a significant impact upon the subduction component in Java and hence the geochemical and isotopic composition of the erupted volcanic rocks. The peak in Ba/Hf and Ba/Nb at Kelut and Tengger Caldera (~112-113°E) (Fig. 7c and inset) corresponds to the centre of the region affected by collision of the Roo Rise (Figs. 1 and 2), and Ba/Hf (and Ba/Nb) decreases relatively systematically at volcanoes either side of this location. The bathymetric survey by Kopp et al. (2006) shows that the outer rise region of the Indian Ocean plate south of Central and East Java is extensively fractured (trench-parallel normal faults) compared to West Java, related to plate bending induced tectonic stress. The faulting and

associated morphological effects caused by subduction of the Roo Rise will increase surface roughness and, therefore, the surface area available for interaction of the subducted lithosphere with seawater (e.g., Ranero et al., 2003). The higher fluid-mobile/non-fluid mobile ratios in East Javanese volcanic rocks may reflect the enhanced water content of the heavily fractured, down-going altered and/or serpentinitised oceanic lithosphere in this region. Enhanced fluid addition and, therefore, fluid-fluxing to the mantle wedge may be expected to result in higher degrees of partial melting of the mantle source. However, the general lack of correlation between Ba/Hf and indicators of the degree of mantle melting (e.g., La/Yb; Fig. 9) in East and Central Javanese volcanic rocks (excluding Ungaran), which show a wide range in Ba/Hf at relatively constant La/Yb, is inconsistent with a systematic increase in fluid-flux melting towards Kelut from the East (Ijen) and West (~Slamet). We do note though that Kelut volcanic rocks have the highest Ba/Hf and amongst the lowest La/Yb ratios of Javanese volcanic rocks, indicative of the greatest degrees of partial melting of the source as might be expected at the centre of the collision-affected region. In contrast, strong correlations between concentrations of fluid-mobile elements (e.g., B), B/La, B/Be and Sr isotope ratios and degrees of mantle melting (La/Yb) have been observed at the Aleutian arc and attributed to subduction of the Amila Fracture Zone in the central Aleutian arc (Singer et al., 1996). Subduction of the fracture zone is proposed to have channelled large quantities of sediment and water into the subarc mantle (Singer et al., 1996). Despite the low La/Yb observed in rocks from Kelut, the overall limited correlation in Javanese rocks of Ba/Hf with La/Yb suggests that the addition of expelled fluid may take place at relatively shallow depth, or there is a greater involvement of high Ba concentration, subducted sedimentary material in volcanic petrogenesis in East and possibly Central Java, relative to West Java. The generally low initial concentration of Ba in serpentinite and experimental serpentinite fluid/residue partition coefficients, led Tenthorey and Hermann (2004) to conclude that Ba enrichment in arc lavas must be largely derived from the subducted sediments. However, the increased fluid-flux available from the heavily-fractured Roo Rise lithosphere likely enables larger degrees of melting of the subducted sediments and scavenging of LILE from the altered oceanic crust (e.g., Ranero et al., 2003). The degree of subducted lithospheric bend faulting, and the incoming plate composition have been linked to the regional trends in lava chemistry for Central American arc volcanism (Rüpke et al., 2002). The transition from subduction accretion (west of ~109°E at the Java Trench) to subduction erosion (east of ~109°E at the Java Trench) (Kopp et al., 2006) may also play a role in providing additional high-Ba pelagic sediments (that were formally part of the accretionary wedge) to the arc element-recycling

budget in East Java. Several studies have suggested that the global subduction erosion flux may be significantly larger than the global subducted sediment flux (e.g., von Huene and Scholl, 1991; Clift et al., 2009; Scholl and von Huene, 2009).

7. Conclusions

New Pb isotope data for Javanese volcanoes and local upper crust, in conjunction with O isotope and geochemical data, have provided new insight into the relative importance of crustal assimilation versus subducted crustal input in Javanese volcanic petrogenesis. Despite varying degrees of modification of geochemical and isotope data by crustal assimilation processes, heterogeneity in the subducted input (fluid and/or sediment melt) is required to explain the significant geochemical and isotopic contrasts observed between West and East Java (e.g., Ba concentration, Ba/Hf ratio and Pb isotopic composition).

We identify two crustal assimilants in West and Central Java: 1) a low $\delta^{18}\text{O}$ and relatively low Pb and Sr isotopic composition assimilant in West Java (at Gede Volcanic Complex), likely representing more primitive arc rocks or mafic-ultramafic ophiolitic rocks, known to outcrop in West (and Central) Java and 2) a higher $\delta^{18}\text{O}$, higher Sr isotope composition assimilant in Central Java, which is consistent with the proposed assimilation of crustal carbonate material for Merapi in Central Java (Chadwick et al., 2007; Deegan et al. 2010; Troll et al., 2013), providing that the carbonate assimilant is characterised by a relatively unradiogenic to moderately radiogenic Pb isotopic composition and low Ba/Hf. However, assimilation of carbonate material in Central Java cannot exert much control the geochemical composition of the erupted rocks in comparison to Vesuvius volcano, where the dominance of carbonate assimilation is clearly documented and manifest in the negative correlation of $\text{Na}_2\text{O}+\text{K}_2\text{O}$ with SiO_2 (Iacono Marziano et al., 2008). Positive correlations between $\text{Na}_2\text{O}+\text{K}_2\text{O}$ with SiO_2 are observed in the majority of Javanese volcanic rocks. The East Javanese volcanic rocks provide little evidence for isotopic modification via crustal assimilation. We suggest that this is due to significant transitions in the crustal architecture and composition between West and Central and Eastern Java at the Central Java Fault/Progo-Muria lineament.

After consideration of the geochemical and isotopic effects of crustal assimilation, the remaining significant heterogeneity observed in East and West Java volcanic rock Ba concentrations, Ba/Hf ratios and Pb isotopic compositions is likely attributed to heterogeneity in the subduction input source component. The high Ba/Hf and lower Pb isotope ratios in East Javanese relative to West Javanese rocks implicate a higher fluid-component and/or greater

incorporation of high Ba/Hf detrital-poor (clay-rich) pelagic sediments in East Java relative to West Java. The collision of the Roo Rise with the Java margin between 109°E and 115°E may provide additional fluid-flux (and greater scavenging of fluid mobile elements) and increased melting of sediment. The transition from subduction accretion to subduction erosion from West to East Java may also play an important role in increasing the budget of crustal material available for subduction recycling in East Java. The tight Javanese Pb isotope array projects back towards a composition consistent with an average I-MORB source.

We note for future studies that: 1) if limestone/carbonaceous sediment assimilation is significant, the ‘least evolved rocks’ of a volcanic suite may not represent the least ‘contaminated’ magma, and 2) where there is a wide variation in isotopic ratios over a limited range in SiO₂ or MgO contents, along-arc studies selecting just one or two basaltic samples to characterise an entire volcano may not accurately represent the most mantle source-like composition. We stress that only through detailed studies of individual volcanoes, prior to along-arc syntheses, will it be possible to fully constrain the relative roles of different crustal inputs in arc settings.

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Figure Captions

Fig. 1. a) Tectonic setting of Java within the Sunda arc, Indonesia. Dashed lines with labels 1-4 represent the Mesozoic and Cenozoic growth of Sundaland suggested by Hall (2011): Line 1: pre-Cretaceous Sundaland Core; Line 2: material added to the pre-Cretaceous Sundaland Core in the Early Cretaceous; Line 3: marks the eastern limit of material added in the Mid Cretaceous; Line 4: marks the eastern boundary of material added in the Early Miocene. The part of Line 2 crossing the Java Sea from Java to Borneo equates to Hamilton’s (1979) SE limit of Cretaceous continental crust. A-D (green fill) represent outcrops of Pre-Tertiary ophiolitic, accretionary-collision complexes in Java and SE Borneo (Hutchison, 1975; Wakita, 2000): A: Ciletuh; B: Luk Ulo (also known as Lok Ulo and Loh Ulo and Karangsambung); C: Jiwo Hills; D: Meratus and Pulau Laut. b) Bathymetric and topographic map of Java and the Java margin compiled from SRTM data. The visible displacement and retreat of the Java Trench by ~60 km to the north between 109°E and 115°E (area between dashed and solid black line (Kopp et al., 2006)) is attributed to the collision and subduction of oceanic basement relief of the Roo Rise (e.g., Kopp et al., 2006). Labels 1-5 represent isolated bathymetric highs that are thought to represent seamounts already subducted at the Java Trench (Kopp et al., 2006). The inferred Progo-Muria lineament (Smyth et al., 2005; Smyth et al., 2007) and Central Java Fault (Chotin et al., 1984; Hoffmann-Rothe et al., 2001) are displayed and discussed in the text. The Eastern Wharton Basin and Argo Basin volcanic provinces (I-Seamount East field in Fig. 3; Hoernle et al., 2011) are delimited by white dashed lines. c) Topographic map of Java showing the location of major volcanoes. Volcanoes for which new Pb and O isotope data are presented in this study along with the location of the three local crustal sedimentary rocks (SED) are marked by red rectangles. Volcano names are consistent with those given by the Smithsonian Institution, Global Volcanism Program (<http://www.volcano.si.edu/>). Division of volcanoes into West, Central and East as defined in the text.

Fig. 2. Along-arc variations for selected Javanese volcanoes showing: a) Subducting seafloor age, b) Average slab dip, c) Slab depth and, d) Distance from trench using data from Syracuse and Abers (2006). Dashed vertical lines denote West, Central and East Java boundaries used

in the manuscript and detailed in Section 3. Arrow shows volcanoes located within the plate margin region affected by the collision and subduction of the Roo Rise (Fig. 1b), an area of oceanic basement relief, as delimited by Kopp et al. (2006). Volcanoes for which geochemical data are used within this study are shown by bold outline symbols and defined in the key.

Fig. 3. a) and b) Pb isotope compositions of Salak, Gede, Galunggung, Merbabu, Merapi and Ijen (this study, colour-filled symbols) compared to published Javanese volcanic rocks (Edwards 1990, 1993; Woodhead et al., 2001), Indian Ocean mid ocean ridge basalt (I-MORB: Rehkämper and Hofmann, 1997; Ito et al., 1987; Price et al., 1986; Chauvel and Blichert-Toft, 2001), Indian Ocean sediments (I-SED: Ben Othman et al., 1989; Gasparon and Varne, 1998), Indian Ocean ferromanganese nodule crusts (I-Mn crusts: Frank and O’Nions, 1998; O’Nions et al., 1998) and Indian Ocean Seamounts (I-Seamount: Outsider Seamount and Cocos-Keeling, Vening-Meinesz, Eastern Wharton Basin and Argo Basin Volcanic Provinces: Hoernle et al., 2011). The composition of seamounts subducting at the Java Trench is represented by the Eastern Wharton Basin and Argo Basin Volcanic Provinces (I-Seamount East). Also plotted are the estimated altered oceanic crust composition (AOC) from Edwards et al. (1993), estimated bulk Java subducted sediment (Plank and Langmuir, 1998), global subducted sediment (GLOSS; Plank and Langmuir, 1998). Local crustal rocks from West Java (marl, volcanoclastic sandstone and mudstone) (this study) along with calcareous sediments from Central Java (Gertisser and Keller, 2003a) are shown. The calc-silicate xenolith is from a Merapi volcanic rock (Gertisser and Keller, 2003a). Sumatran intrusive rocks are from Gasparon and Varne (1995). An example bulk mixing line (0-100%) is shown between average I-MORB source (Pb: 0.07 ppm, $^{206}\text{Pb}/^{204}\text{Pb}$: 17.899, $^{207}\text{Pb}/^{204}\text{Pb}$: 15.471, $^{208}\text{Pb}/^{204}\text{Pb}$: 37.763, n = 43) and bulk Java subducting sediment (Pb: 25.5 ppm, $^{206}\text{Pb}/^{204}\text{Pb}$: 18.990, $^{207}\text{Pb}/^{204}\text{Pb}$: 15.741, $^{208}\text{Pb}/^{204}\text{Pb}$: 39.328, Plank and Langmuir, 1998). c) and d) show the new Pb isotope data for Salak, Gede, Galunggung, Merbabu, Merapi and Ijen volcanic rocks (large symbols) and new local crust data from West Java (marl, volcanoclastic sandstone and mudstone) with other published Javanese (including Krakatau) volcano Pb isotope data (small symbols; Edwards, 1990, 1993; Woodhead et al., 2001).

Fig. 4. a) Olivine (ol), clinopyroxene (cpx) and plagioclase (plag) mineral separate $\delta^{18}\text{O}$ and whole-rock (WR) (inset) $\delta^{18}\text{O}$ values versus $^{206}\text{Pb}/^{204}\text{Pb}$ for Javanese volcanic rocks. Whole-rock Merapi Pb isotope data is from this study and the O isotope data is from Gertisser and

Keller (2003a). Mineral data are plotted as measured $\delta^{18}\text{O}$ values, rather than calculated equilibrium melt values, due to the potential uncertainty in mineral-melt fractionation factors compared to mineral-mineral fractionation factors (Eiler, 2001). b) Along-arc variations in $\delta^{18}\text{O}$ of Javanese volcano mineral separates and whole-rock samples (inset). The Merbabu volcanic rocks are amongst the highest $\delta^{18}\text{O}$ whole-rock values yet reported in Java. S: Salak (Handley et al., 2008a), Ge: Gede (Handley et al., 2010), Gu: Guntur (Edwards, 1990; Macpherson et al., submitted), Ga: Galunggung (Gerbe et al., 1992), C: Cereme (Edwards, 1990), M: Merapi (Gertisser and Keller, 2003a; Troll et al., 2013), K: Kelut (Jeffery et al., 2013), I: Ijen (Handley et al., 2007). The upper mantle range of $+5.5 \pm 0.2\text{‰}$ for clinopyroxene is taken from Eiler (2001). c) Javanese clinopyroxene mineral separate $\delta^{18}\text{O}$ values versus whole-rock $^{206}\text{Pb}/^{204}\text{Pb}$. Simple bulk-mixing curves are shown for mixing between depleted mantle source and local Indian Ocean sediment (model A) and Javanese arc magma with local arc crust (model B). Tick marks indicate the percentage of crustal material added to the depleted mantle source and arc magma. Data used in mixing calculations: Depleted mantle source: Pb = 0.07 ppm; $^{206}\text{Pb}/^{204}\text{Pb} = 17.899$ (average I-MORB composition used in Fig. 3); O = 43.8 wt% (Vroon et al., 2001); $\delta^{18}\text{O} = +5.5\text{‰}$ (Eiler, 2001). Local subducted sediment: Pb = 25.5 ppm; $^{206}\text{Pb}/^{204}\text{Pb} = 18.990$ (Plank and Langmuir, 1998); O = 50.2 wt% (Vroon et al., 2001); $\delta^{18}\text{O} = 18.7\text{‰}$ (Vroon et al., 2001; based on DSDP site 262 data). Arc magma: Pb = 6.1 ppm (KI 69, Handley et al., 2007); $^{206}\text{Pb}/^{204}\text{Pb} = (\text{KI 69, Table 1})$; O = 50.2 wt% (Vroon et al., 2001); $\delta^{18}\text{O} = +5.5\text{‰}$. Local arc crust: Pb = 4.95 ppm (SEDA, Table 3); $^{206}\text{Pb}/^{204}\text{Pb} = 18.722$ (SEDA, Table 1); O = 50.2 wt% (Vroon et al., 2001); $\delta^{18}\text{O} = 19\text{‰}$ (representative of local calcareous sedimentary arc crust given by Gertisser and Keller, 2003a). Inset diagram shows the expected mixing trends for crustal assimilation (dashed lines) versus subducted sediment input (solid lines) for crustal materials with variable $^{206}\text{Pb}/^{204}\text{Pb}$ ratios. DM = depleted mantle source.

Fig. 5. Variation of whole-rock Pb isotope ratios with SiO_2 content for Salak, Gede, Galunggung, Merapi, Merbabu and Ijen volcanic rocks relative to other Javanese volcanic rocks (Edwards 1990, 1993; Woodhead et al., 2001). SiO_2 data, for the samples with new Pb isotope data, are from Sitorus (1990), Turner and Foden (2001), Gertisser and Keller (2003a), Handley et al. (2007; 2008a; 2010; 2011), Preece et al. (2013) plus ME07-53: 55.40 wt%, M11-05: 54.08 wt%, M11-18: 55.72 wt%. The central Java upper crustal calcareous sedimentary rocks of Gertisser and Keller (2003a) are plotted on the Y-axis (SiO_2 contents not

available). Arrows labelled SH, AFC and FC indicate the hypothesised data trends related to: heterogeneity in the mantle source (SH), combined assimilation and fractional crystallisation (AFC) and fractional crystallisation (FC). AFC trends can be positive or negative depending on the Pb isotope ratio of the assimilated material. AFC 1: high Pb isotope ratio assimilant e.g., continental crust; AFC 2: assimilation of local West and Central Javanese carbonaceous sedimentary rocks; AFC 3: assimilation of more isotopically primitive arc rocks or ophiolitic-type arc crust.

Fig. 6. $\text{Na}_2\text{O}+\text{K}_2\text{O}$ versus SiO_2 for Javanese volcanic rocks and local, West Java upper crustal rocks (marl, volcanoclastic sandstone and mudstone). Large symbols show the volcanoes with new Pb isotope data presented in this study. Arrows (upper left) show the expected differentiation path for fractional crystallisation of clinopyroxene (FC Cpx) and olivine (FC Ol) and indicate the desilication effect of carbonate assimilation in a closed system depending on the availability (Carb Assim (Ol/MgO)) or not (Carb Assim (no Ol)) of MgO in the magma, following Iacono Marziano et al. (2008). Java data sources as in Figs. 3 and 5 plus Claproth (1989), Camus et al. (1987), Gerbe et al. (1992), Vukadinovic and Sutawidjaja (1995), Hartono (1996), van Gerven and Pichler (1995), Mandeville et al. (1996), Carn and Pyle (2001), Jeffery et al. (2013) and Abdurrachman and Yamamoto (2012).

Fig. 7. Along-arc a) Pb isotope, b) Sr isotope and, c) Ba/Hf and Ba/Nb (inset) variations in Javanese volcanic rocks. Note the similarity between the along-arc O isotope (Fig. 4b, clinopyroxene or plagioclase mineral data) and Sr isotope patterns compared to the along-arc Pb isotope pattern. Data sources given in Figs. 3, 5, 6 plus additional data from Whiford (1975) and Abdurrachman and Yamamoto (2012). Da: Danau Complex; TP: Tangkubanparahu; P: Papandayan; Gu: Guntur; Ci: Cikuray; Ga: Galunggung; C: Cereme; Sl: Slamet; D: Dieng Volcanic Complex; Sun: Sundoro; Su: Sumbing; Un: Ungaran; Law: Lawu; Ke: Kelut; Se: Semeru; Lam: Lamongan. Hf concentration data is not available for Tengger Caldera but this volcano is plotted in the East Java field (112.95°E) in the insert Ba/Nb diagram (van Gerven and Pichler, 1995).

Fig. 8. a) Ba concentration versus SiO_2 content in Javanese volcanic rocks. Large symbols show the volcanoes with new Pb isotope data presented in this study. Data sources are given in Figs. 3 and 7. b) Pb isotope ratio versus Ba/Hf versus in Javanese volcanic rocks. Data sources are given in Figs. 3 and 7. Symbols as those given in Fig. 3. Local Indian Ocean

sediment (I-SED) end members used in bulk-mixing models: A: Lower Cretaceous claystone (29-2:29-31), Pb: 15 ppm, $^{206}\text{Pb}/^{204}\text{Pb}$: 18.576, Ba: 1063 ppm, Hf: 2.69 ppm (Gasparon and Varne, 1998); B: Quaternary terrigenous mud (VM33-79) Pb: 17 ppm, $^{206}\text{Pb}/^{204}\text{Pb}$: 18.67, Ba: 129 ppm, Hf: 3.04 ppm (Gasparon and Varne, 1998); C: Bulk Java Sediment, Pb: 25.5 ppm, $^{206}\text{Pb}/^{204}\text{Pb}$: 18.99, Ba: 1068 ppm, Hf: 4.73 ppm (Plank and Langmuir, 1998). Depleted I-MORB mantle source: Pb: 0.07 ppm, $^{206}\text{Pb}/^{204}\text{Pb}$: 17.899, Ba: 2.74 ppm, Hf: 0.25 ppm (Rehkämper and Hofmann, 1997; Ito et al., 1987; Price et al., 1986; Chauvel and Blichert-Toft, 2001). Tick marks show the percentage of bulk sediment added (increments are 0.5%, 1%, 1.5%, 2%, 2.5%, 3%, 5%, 10%, 50%, 100%) to the depleted I-MORB source. Box with arrows (lower right) exemplifies the anticipated maximum overprint on Ba/Hf and $^{206}\text{Pb}/^{204}\text{Pb}$ from crustal assimilation at Gede Volcanic Complex, West Java.

Fig. 9. Ba/Hf versus La/Yb in Javanese volcanic rocks. Data sources are given in Fig. 7.

Table 1. Pb isotope data of Javanese volcanic and sedimentary rocks

Province	Volcano	Eruptive Vent/Age	Sample	$^{206}\text{Pb}/^{204}\text{Pb}$	$^{207}\text{Pb}/^{204}\text{Pb}$	$^{208}\text{Pb}/^{204}\text{Pb}$
West Java	Salak	Central Vent	S109	18.827	15.703	39.222
West Java	Salak	Central Vent	S110A	18.799	15.676	39.150
West Java	Salak	Central Vent	S111	18.814	15.700	39.199
West Java	Salak	Side Vent	S103	18.703	15.670	39.018
West Java	Salak	Side Vent	S107B	18.832	15.704	39.222
West Java	Salak	Pre-Salak	S100	18.816	15.697	39.187
West Java	Gede Volcanic Complex	Young Gede (KR)	G23	18.879	15.715	39.281
West Java	Gede Volcanic Complex	Young Gede (KR)	G25	18.830	15.688	39.155
West Java	Gede Volcanic Complex	Young Gede (KR)	G40	18.870	15.710	39.262
West Java	Gede Volcanic Complex	Young Gede (OV)	G20	18.867	15.707	39.270
West Java	Gede Volcanic Complex	Young Gede (OV)	G36A	18.863	15.706	39.236
West Java	Gede Volcanic Complex	Old Gede	G17	18.837	15.699	39.194
West Java	Gede Volcanic Complex	Old Gede	G46	18.873	15.708	39.262
West Java	Gede Volcanic Complex	Pangrango	G35	18.938	15.733	39.400
West Java	Gede Volcanic Complex	Gegerbentang	G33	18.914	15.725	39.352
West Java	Gede Volcanic Complex	Older Quaternary	G52	18.774	15.687	39.117
West Java	Sedimenatry Rock	Marl	SEDA	18.722	15.665	38.944
West Java	Sedimenatry Rock	Volcanic Sandstone	SEDB	18.687	15.663	38.968
West Java	Sedimenatry Rock	Mudstone	SEDC	18.785	15.675	38.971
West Java	Galunggung*	AD 1982 eruption	VB82	18.7991	15.6987	39.1939
West Java	Galunggung*	AD 1918 eruption	4AK	18.7524	15.6807	39.1007
Central Java	Merapi	AD 2006 eruption	ME07-53	18.762	15.693	39.147
Central Java	Merapi	AD 2006 eruption	ME08-07	18.766	15.697	39.157
Central Java	Merapi	AD 2006 eruption	ME08-14	18.762	15.692	39.143
Central Java	Merapi	AD 2010 eruption	M11-05	18.770	15.696	39.162
Central Java	Merapi	AD 2010 eruption	M11-12	18.762	15.694	39.147
Central Java	Merapi	AD 2010 eruption	M11-18	18.762	15.692	39.125
Central Java	Merapi	AD 2010 eruption	M11-27-5	18.760	15.692	39.137
Central Java	Merapi	AD 2010 eruption	M11-28b	18.765	15.697	39.153
Central Java	Merapi*	High-K, Recent-Historical	M97-068	18.763	15.694	39.141
Central Java	Merapi*	High-K, Holocene	M98-096	18.771	15.697	39.156
Central Java	Merapi*	Med-K, Holocene	M96-137	18.759	15.695	39.139
Central Java	Merapi*	Med-K, Holocene	M96-073	18.755	15.692	39.134
Central Java	Merapi*	Med-K, Merapi-Somma	M95-026	18.769	15.693	39.130
Central Java	Merbabu	S-sector (Patran)	MB-1	18.828	15.710	39.245
Central Java	Merbabu	S-sector (Jrakah-Selo)	MB-2	18.802	15.690	39.173
Central Java	Merbabu	SW-sector (SE of Candimulyo)	MB-6	18.805	15.691	39.182
Central Java	Merbabu	N-sector (E of Kopeng)	MB-16	18.756	15.665	39.040
Central Java	Merbabu	N-sector (NE of Getasan)	MB-22	18.755	15.669	39.045
Central Java	Merbabu	E-sector (N or Penggung)	MB-28	18.807	15.696	39.188
East Java	Ijen Volcanic Complex	Djampit (Caldera Rim)	KI 69	18.526	15.617	38.768
East Java	Ijen Volcanic Complex	Djampit (Caldera Rim)	KI 137	18.551	15.611	38.753
East Java	Ijen Volcanic Complex	Djampit (Caldera Rim)	KI 136	18.540	15.611	38.747
East Java	Ijen Volcanic Complex	Merapi (Caldera Rim)	KI 116	18.578	15.626	38.802
East Java	Ijen Volcanic Complex	Merapi (Caldera Rim)	KI 194	18.598	15.628	38.836
East Java	Ijen Volcanic Complex	Kawah Ijen	KI 190	18.602	15.635	38.861
East Java	Ijen Volcanic Complex	Glaman (Intra Caldera)	KI 35	18.574	15.624	38.806
East Java	Ijen Volcanic Complex	Anyar (Intra Caldera)	KI 142	18.563	15.623	38.789

Further sample information is available in Handley et al. (2007) for Ijen, Handley et al. (2008) for Salak, Gertisser et al. (2003a, 2003b) and Preece et al. (2013) for merapi, Handley et al. (2011) for Merbabu and Handley et al. (2010) for Gede Volcanic Complex.

KR: Kawa Ratu; OV: Other Vents Group

*samples re-analysed in this study for which previously published TIMS Pb isotope data is available in Gertisser et al. (2003a) for Merapi and Turner et al. (2001) for Galunggung.

Data comparison between the new and previously published Pb isotope data is provided in the Appendix.

Table 2. Whole-rock oxygen isotope compositions of Merbabu volcanic rocks

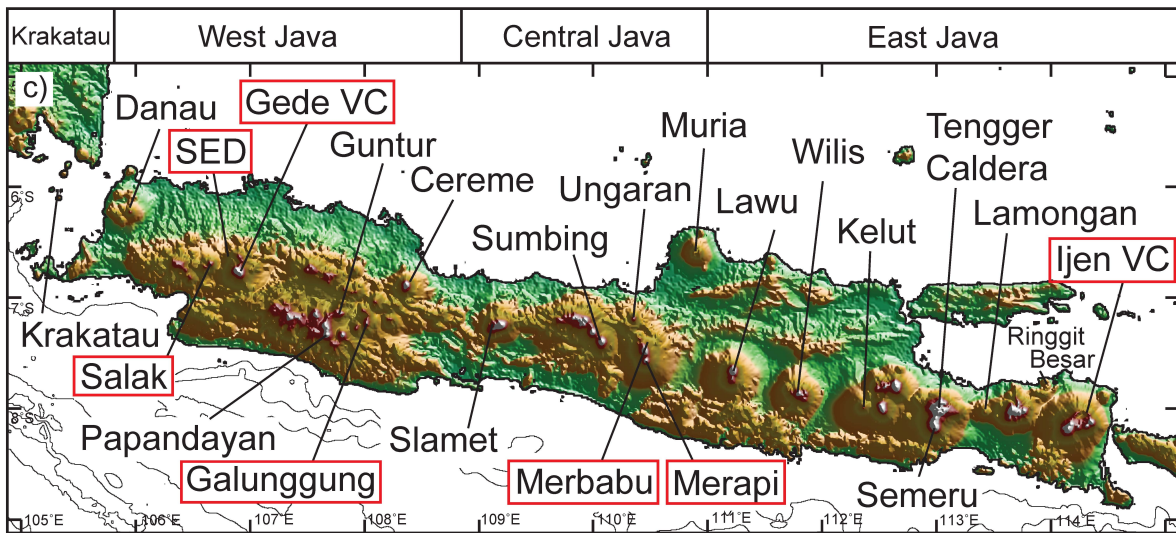
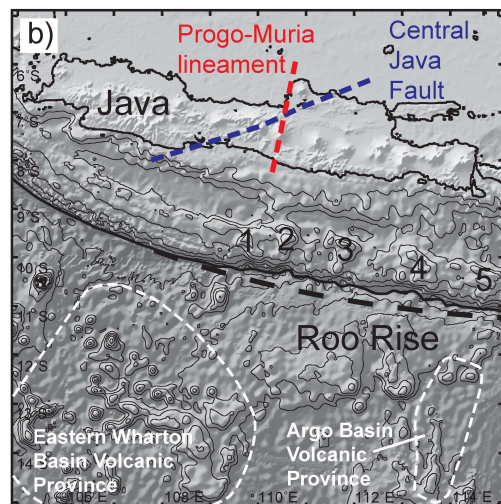
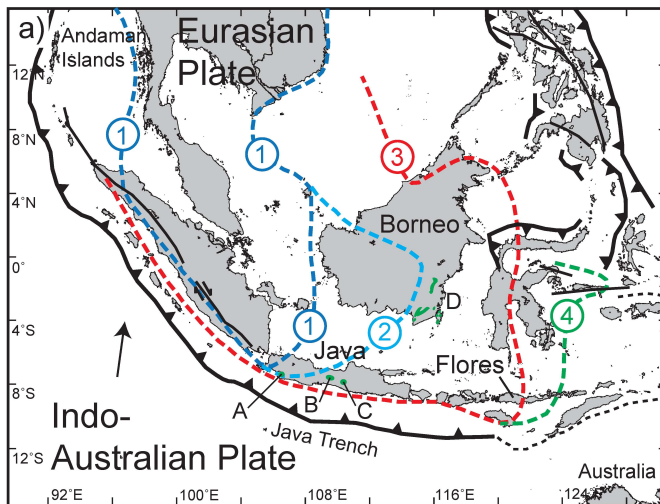
Sample	$\delta^{18}\text{O}$
MB-1	+7.3
MB-2	+6.4
MB-6	+7.2
MB-16	+8.4
MB-22	+8.1
MB-28	+6.8

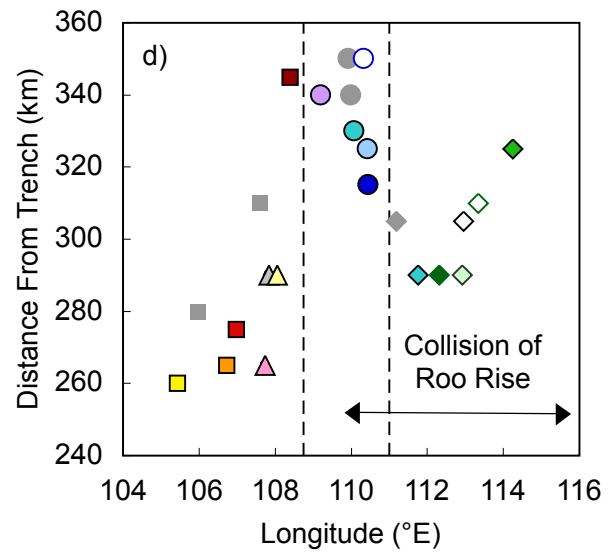
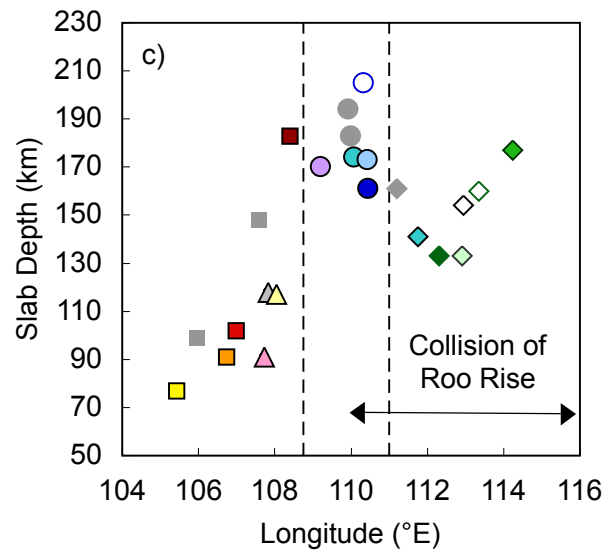
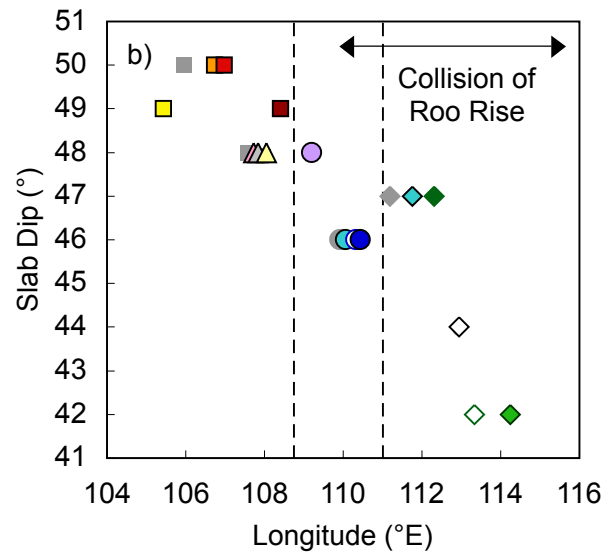
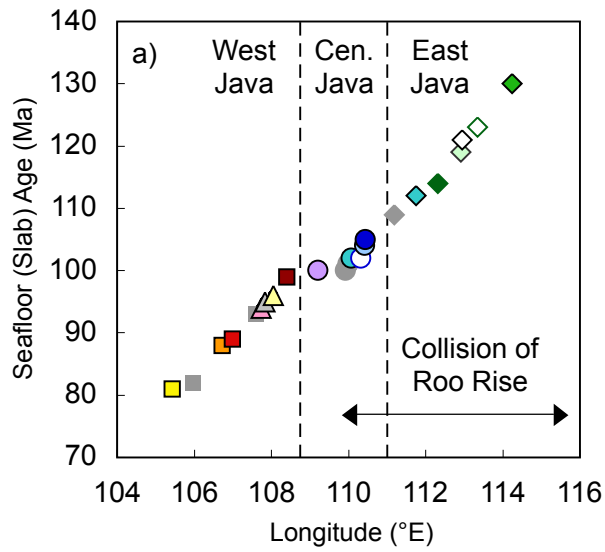
O isotope data reported as per mil (‰)

Table 3. Major element, trace element and Sr-Nd-Hf isotope data of West Java upper crust sedimentary rocks

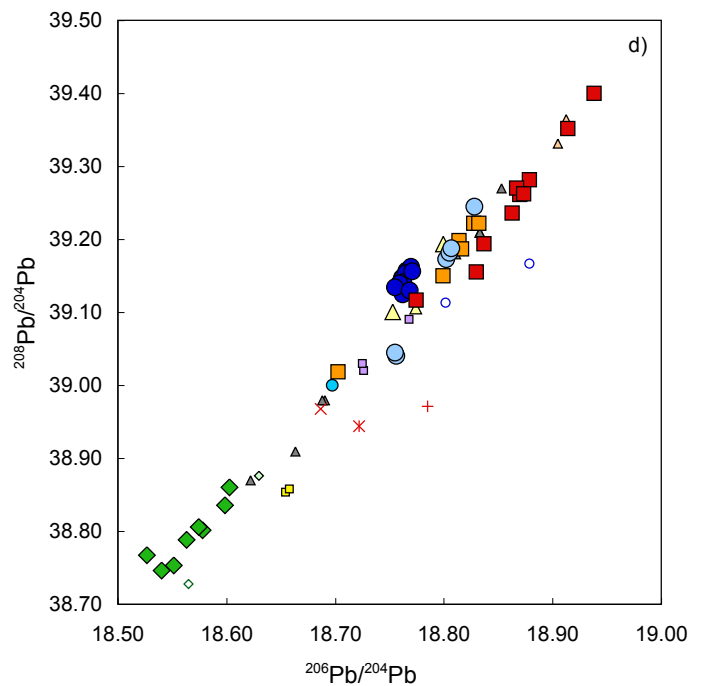
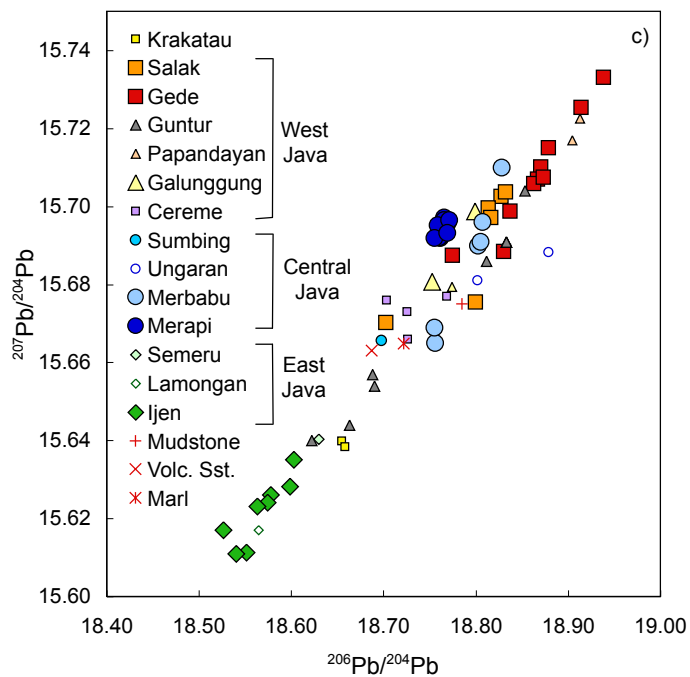
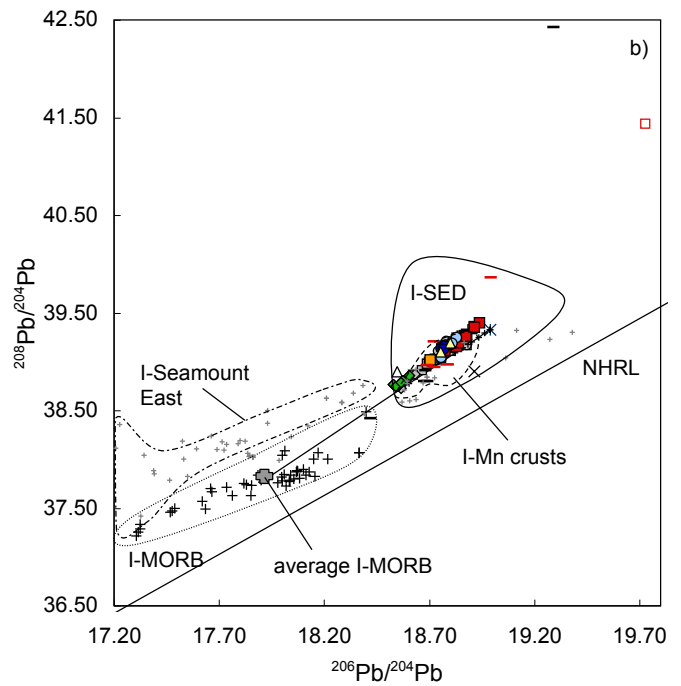
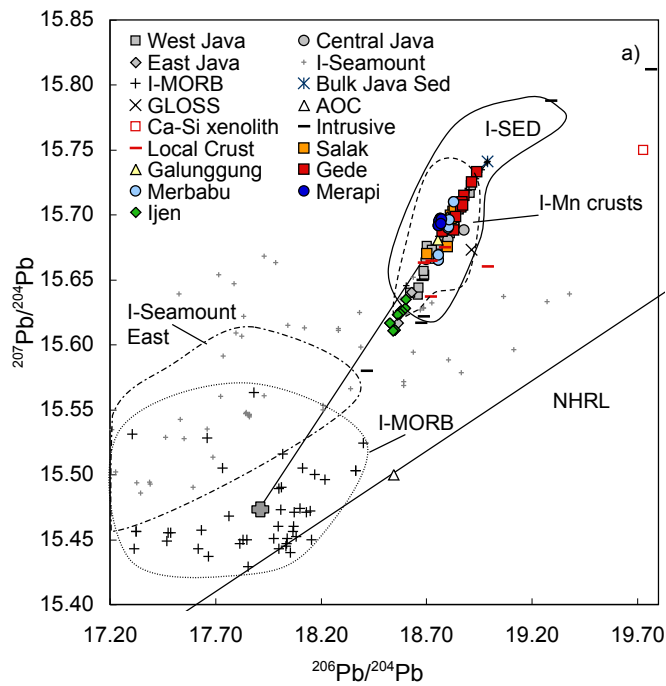
Sample	SED-A	SED-B	SED-C
Latitude	06°37'30.7"S	06°37'30.7"S	06°37'30.7"S
Longitude	106°53'06.4"E	106°53'06.4"E	106°53'06.4"E
Elevation	438 ± 7m	438 ± 7m	438 ± 7m
Rock Type	Marl	Volcaniclastic sandstone	Mudstone
SiO ₂	14.99	51.42	55.74
Al ₂ O ₃	5.61	20.67	22.29
Fe ₂ O ₃	4.94	6.61	6.53
MgO	8.63	2.10	2.13
CaO	28.71	6.73	0.72
Na ₂ O	0.55	2.93	0.36
K ₂ O	0.34	0.65	2.01
TiO ₂	0.21	0.78	0.87
MnO	1.64	0.11	0.05
P ₂ O ₅	0.14	0.17	0.05
LOI	32.97	6.47	9.0
Total	98.72	98.64	99.75
Na ₂ O+K ₂ O	0.886	3.576	2.367
Sc	2.5	12.8	17.0
V	33	34	118
Cr	7.7	3.0	47.8
Co	3.5	8.0	14.1
Ni	7.5	4.6	23.5
Cu	5.6	11.3	14.6
Zn	27.6	76.6	88.4
Rb	9.0	10.0	76.4
Sr	373	372	74
Y	29.6	24.1	20.1
Zr	66	186	165
Nb	3.4	15.4	10.2
Cs	0.5	0.2	6.8
Ba	73.8	141.6	155.7
La	15.8	27.3	27.7
Ce	30.3	61.2	62.3
Pr	3.84	8.05	7.15
Nd	16.2	32.3	27.7
Sm	3.45	6.20	5.35
Eu	0.83	2.06	1.14
Gd	3.66	4.98	4.42
Tb	0.60	0.81	0.69
Dy	3.80	4.69	3.93
Ho	0.85	0.95	0.80
Er	2.55	2.57	2.26
Tm	0.43	0.41	0.37
Yb	2.87	2.70	2.41
Lu	0.51	0.45	0.40
Hf	1.46	5.05	4.01
Ta	0.26	0.99	0.83
Pb (total)	4.95	13.6	23.9
Th	2.93	7.31	12.4
U	1.91	1.73	2.21
Ba/Hf	50.7	28.1	38.8
⁸⁷ Sr/ ⁸⁶ Sr	0.704339	0.704979	0.709587
2SE	0.000008	0.000011	0.000011
¹⁴³ Nd/ ¹⁴⁴ Nd	0.512806	0.512792	0.512552
2SE	0.000032	0.000007	0.000006
¹⁷⁶ Hf/ ¹⁷⁷ Hf	0.283313	0.283077	0.282966
2SE	0.000082	0.000007	0.000007

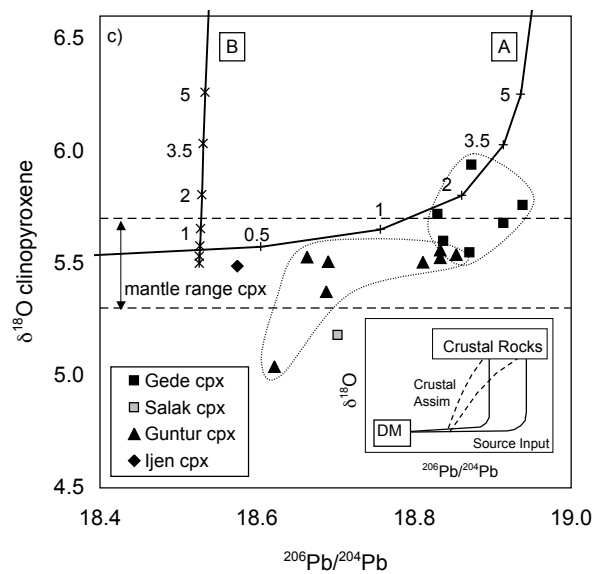
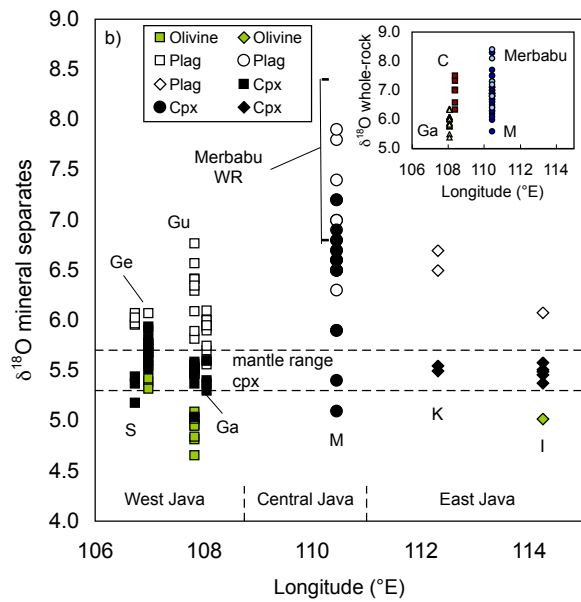
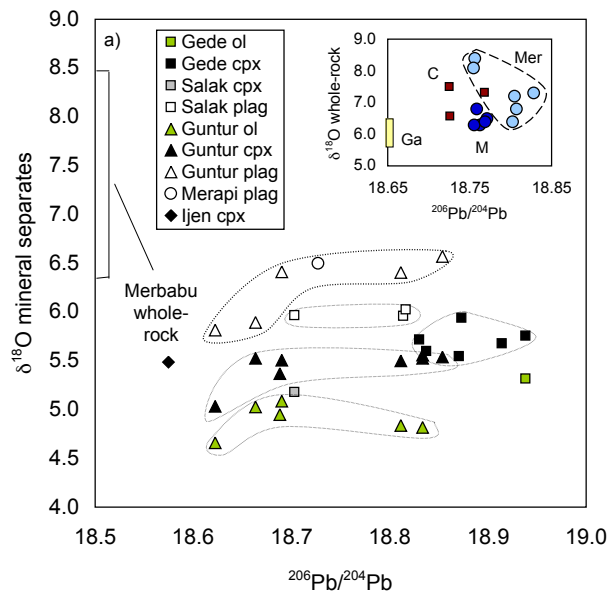
Major element contents are given in wt% and trace element concentrations are given in ppm.

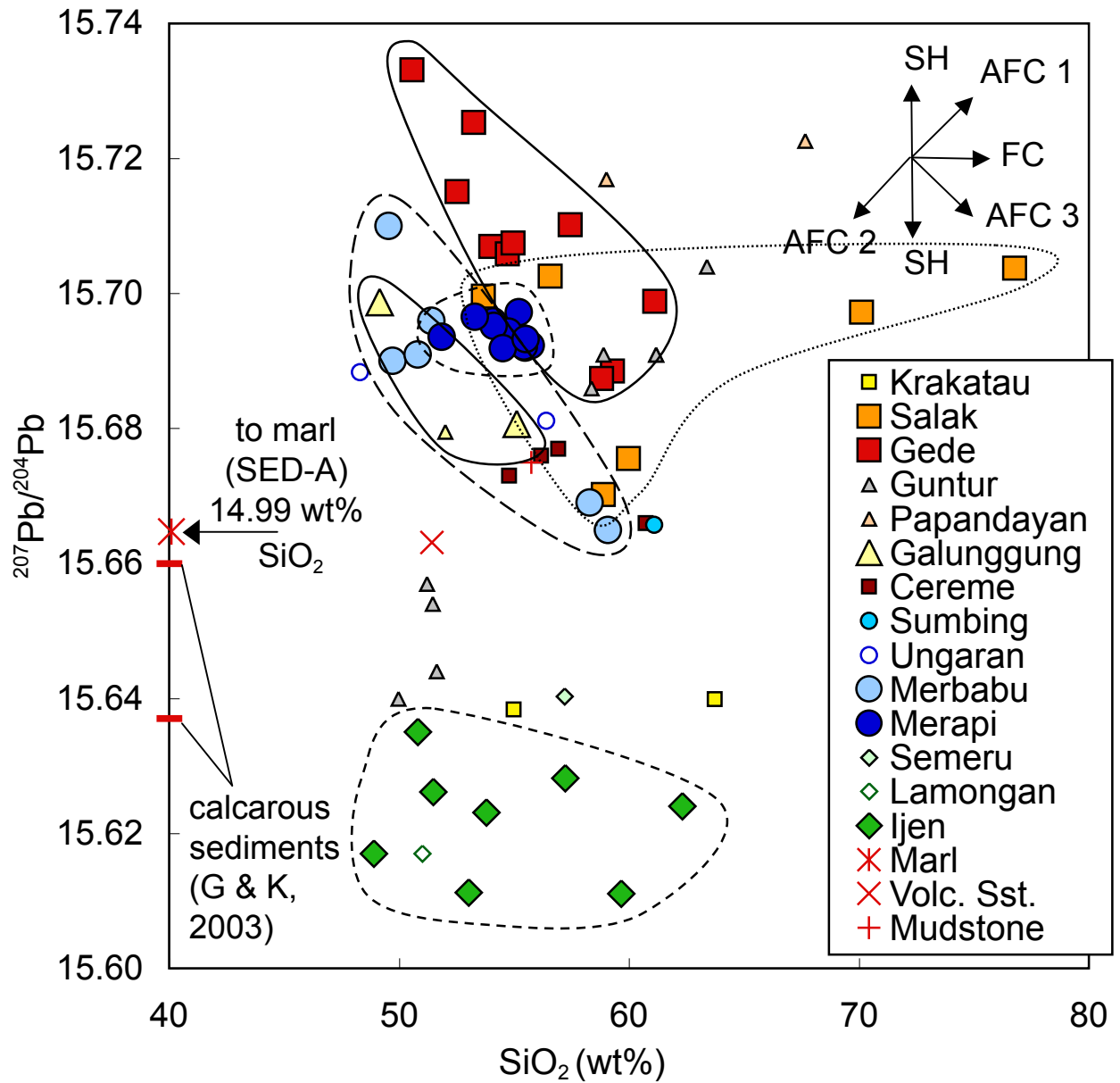


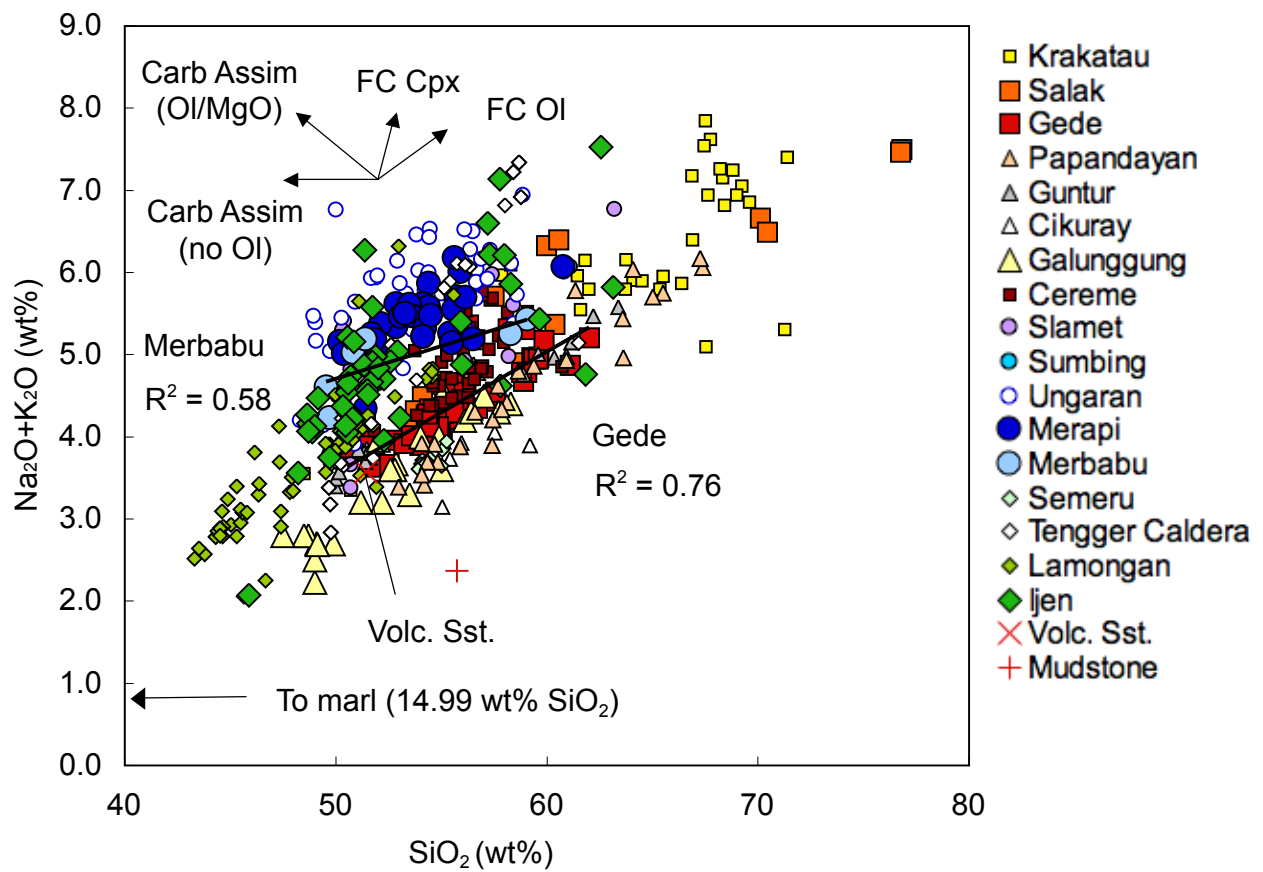


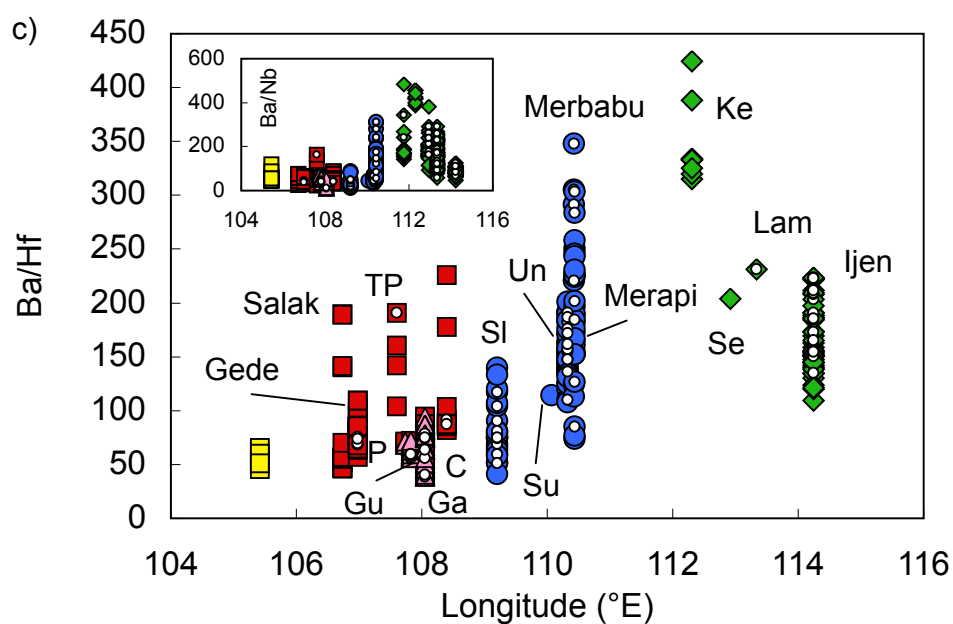
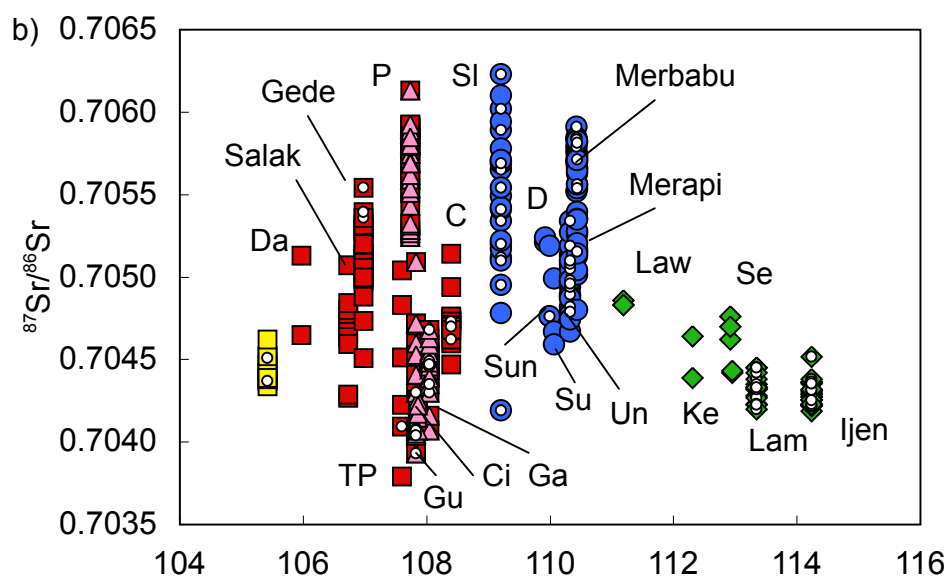
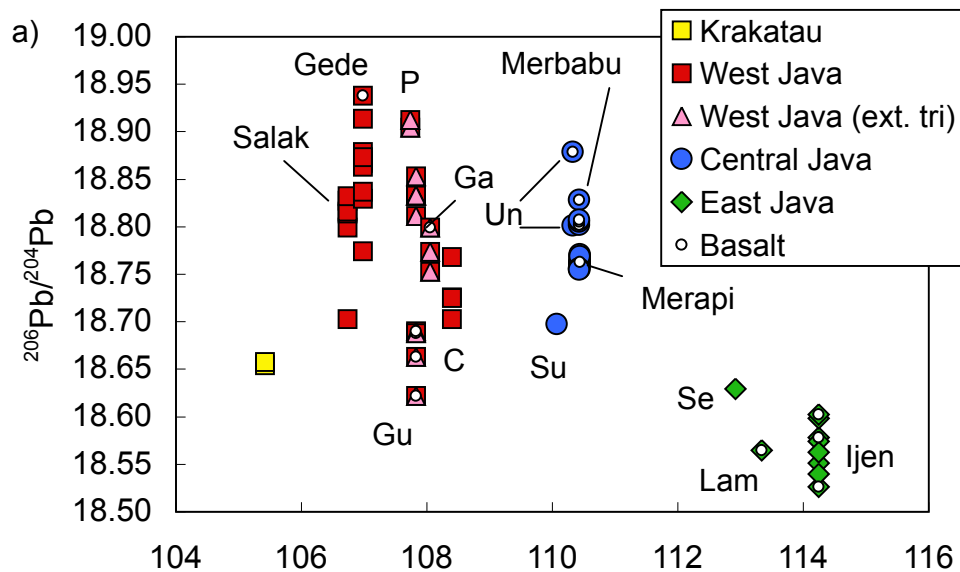
- | | |
|--------------|-------------------|
| ■ Krakatau | ■ Salak |
| ■ Gede | △ Guntur |
| △ Papandayan | △ Cikuray |
| △ Galunggung | ■ Cereme |
| ● Slamet | ● Sumbing |
| ○ Ungaran | ● Merapi |
| ● Merbabu | ◆ Wilis |
| ◆ Semeru | ◆ Lamongan |
| ◆ Ijen | ◇ Tengger Caldera |
| ◆ Kelut | |

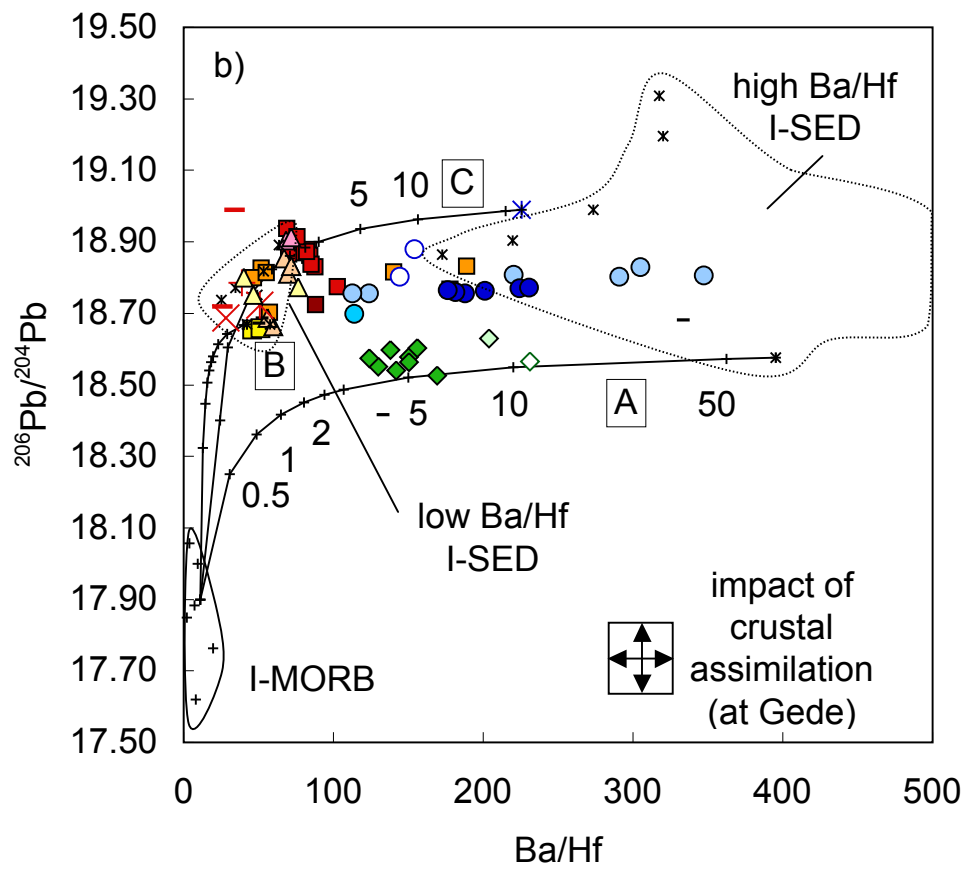
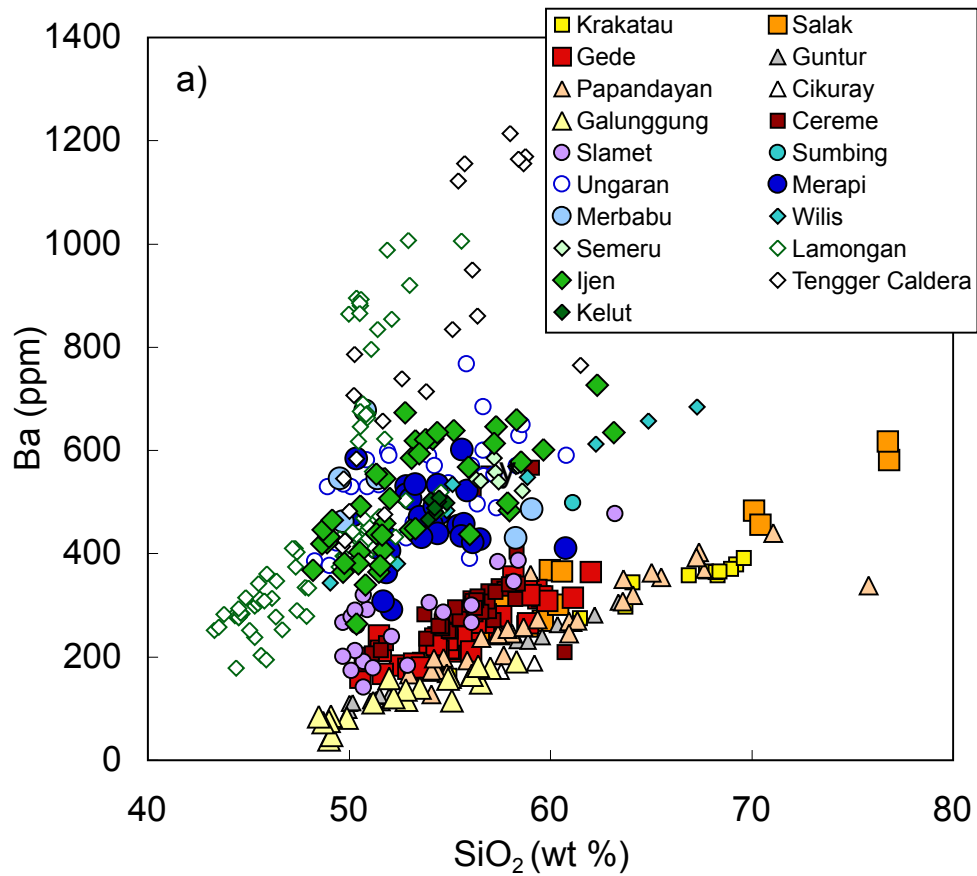


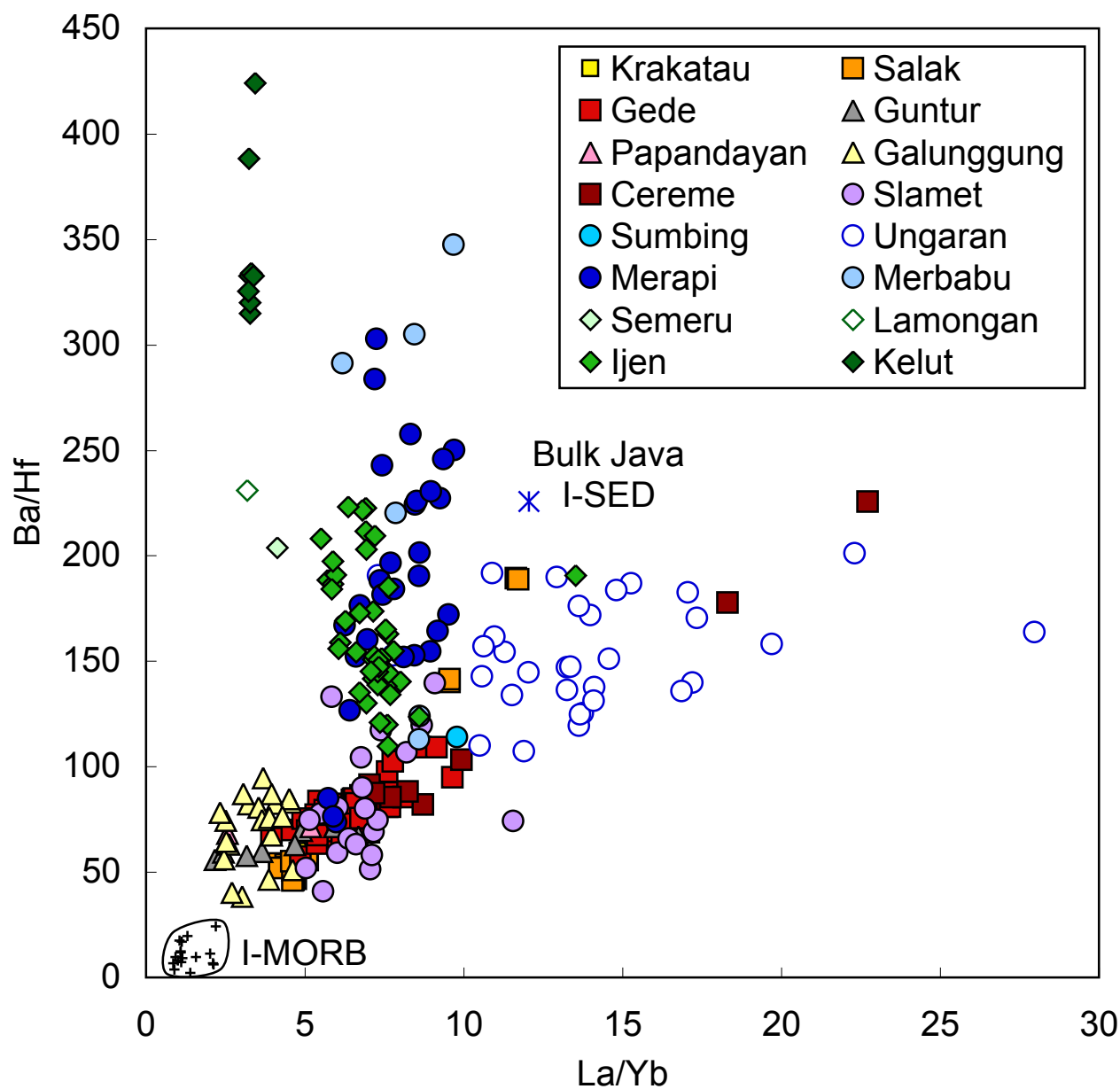












Appendix

Comparison of re-analysed Pb isotope ratios with previously published data

An inter-laboratory comparison of Pb isotope data by Thirlwall (2000) revealed significant variation in reported Pb isotope ratios for the same samples (up to 30 times the supposed reproducibility based upon replicate analyses of NIST SRM 981). These large differences were attributed to inadequate fractionation corrections and/or environmental contamination of rock powder. Woodhead and Hergt (2000) also showed that the conventional corrections for (unspiked) thermal ionisation mass spectrometer (TIMS) induced Pb-isotopic fractionation often result in a loss of accuracy due to the different fractionation behaviours of pure Pb reference materials (NIST SRM 981, 982) and natural rock samples. Therefore, rock powders from Galunggung and Merapi volcanoes with previously published Pb isotope data were re-analysed in this study to facilitate comparison between data sets collected in different laboratories using different methods and instrumentation (e.g., Tl-doped multi-collector inductively-coupled plasma mass spectrometry (MC-ICP-MS) with sample-standard bracketing data versus unspiked, conventionally fractionation-corrected TIMS data).

Table A1 compares the analytical techniques employed for the previously published (TIMS) and new (re-analysed) Pb isotope data for Galunggung and Merapi. The new and previously published Pb isotope data are presented in Table A2 and Figure A1.

Table A1. Summary of analytical methods for new (re-analysed) and previously published data for Galunggung and Merapi volcanoes

Volcano	Data Source	Instrument and Laboratory	Pre-sample Treatment	Fractionation Correction Control
Galunggung	Turner and Foden (2001)	TIMS (Open University)	None	~1‰ per atomic mass unit using NIST SRM 981 values from Todt et al. (1993)
Merapi	Gertisser and Keller (2003a)	TIMS (Universität Tübingen)	None	1‰ per atomic mass unit using NIST SRM 981 (value not given)
Galunggung and Merapi	This study	MC-ICP-MS	Leached (hot 6 M HCl)	Tl doping and sample-standard bracketing using NIST SRM 981 and values from Eisele et al. (2003)

Table A2. Comparison of new (re-analysed) and previously published Pb isotope data of Galunggung and Merapi volcanic rocks

Volcano/ Sample	Data source	$^{206}\text{Pb}/^{204}\text{Pb}$	$^{207}\text{Pb}/^{204}\text{Pb}$	$^{208}\text{Pb}/^{204}\text{Pb}$
Galunggung				
VB82	Turner and Foden (2001)	18.735	15.627	38.947
4AK	Turner and Foden (2001)	18.692	15.624	38.862
VB82	This study	18.799	15.699	39.194
4AK	This study	18.752	15.681	39.101
Merapi				
M97-68	Gertisser and Keller (2003a)	18.727	15.671	39.038
M96-137	Gertisser and Keller (2003a)	18.727	15.681	39.015
M96-73	Gertisser and Keller (2003a)	18.757	15.695	39.138
M98-96	Gertisser and Keller (2003a)	18.775	15.702	39.178
M95-26	Gertisser and Keller (2003a)	18.784	15.716	39.208
M97-68	This study	18.763	15.694	39.141
M96-137	This study	18.759	15.695	39.139
M96-73	This study	18.755	15.692	39.134
M98-96	This study	18.771	15.697	39.156
M95-26	This study	18.769	15.693	39.130

The previously published and new, re-analysed data for the same rock powders from Galunggung (VB82 and 4AK) show considerable variation in measured Pb isotope ratios (Table A2 and Fig. A1). The Galunggung Pb isotope ratios determined by Turner and Foden (2001) plot at significantly lower $^{207}\text{Pb}/^{204}\text{Pb}$ and slightly lower $^{206}\text{Pb}/^{204}\text{Pb}$ relative to both the new re-analysed data and the main Java array. The re-analysed Merapi samples show more similar Pb isotopic compositions to the previously published TIMS data of Gertisser and Keller (2003a) but the re-analysed data cover a more restricted range, particularly for $^{207}\text{Pb}/^{204}\text{Pb}$. The differences between the previously published TIMS data and our re-analysed MC-ICP-MS data for Galunngung and Merapi are likely due to: 1) the pre-treatment leaching in HCl for the re-analysed samples versus no leaching of the samples analysed by Turner and Foden (2001) and Gertisser and Keller (2003a) and 2) the conventional corrections employed for (unspiked) thermal ionisation mass spectrometer (TIMS) induced Pb-isotopic fractionation for the previously published data, versus Tl-doping and MC-ICP-MS sample-standard bracketing for the re-analysed samples.

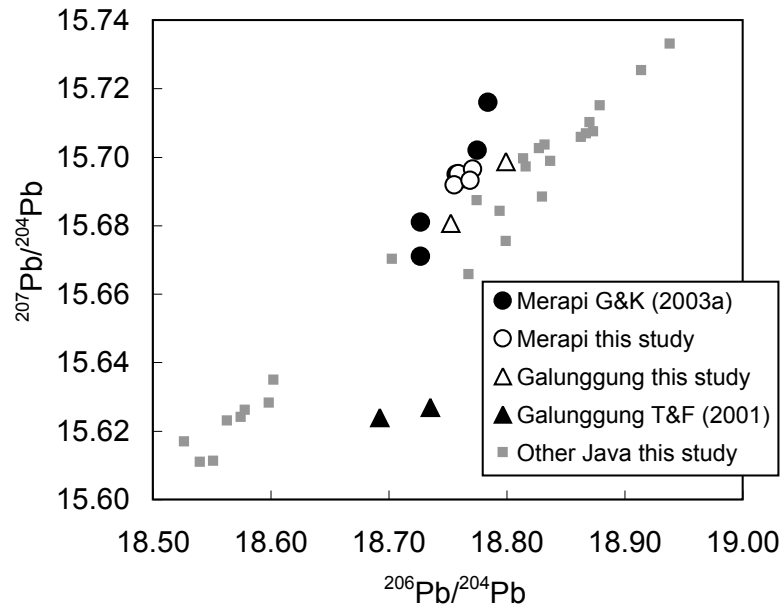


Fig. A1. Comparison of re-analysed and previously published Pb isotope data of Galunggung (Turner and Foden, 2001) and Merapi (Gertisser and Keller, 2003a) volcanic rocks. The new data for Gede, Salak and Ijen volcanoes (this study) are also plotted (grey symbols) for reference to the main Java array.

Selection of Javanese Pb isotope data sets for comparison

Due to the significant discrepancy observed between the previously published and re-analysed Galunggung Pb isotope data we have not used the Javanese Pb isotope data of Turner and Foden (2001) as a comparison data set in the manuscript. We use our re-analysed Pb isotope data for Merapi (instead of the Pb isotope data published by Gertisser and Keller, 2003a) in order for the data to be directly comparable to the new Merapi data presented in this study for the volcanic rocks erupted in 2006 and 2010. We note that as the Merbabu samples in this study were analysed using the same analytical methods and instrumentation of Gertisser and Keller (2003a) these data may also show a greater spread in $^{207}\text{Pb}/^{206}\text{Pb}$ relative to the new data presented for Merapi. Therefore, in the manuscript no attention has been drawn to the fact that the Merbabu Pb isotope data define an array of slightly different slope compared to the overall slope of the main Java array. The previously published Merapi data also presented a similar slope to Merbabu (compare the Merapi data of Fig. A2 with Merbabu data of Fig. 3c in the manuscript). The two Javanese data sets utilised for comparison in this study are Woodhead et al. (2001) (double-spiked TIMS data) and Edwards (1990) (conventional fractionation corrected TIMS data). Despite the latter study using a conventional fractionation correction, the Cereme and Guntur

data of Edwards (1990) plot along the main Java array and show the same slope (Fig. 3 of the manuscript). The older Javanese Pb isotope data set of Whitford (1975) has also been excluded from data comparison due to similar and significant discrepancies visible between the Whitford (1975) data and samples that were re-analysed by Woodhead et al. (2001).

Extended analytical techniques for the Tertiary sedimentary rocks of West Java

Major element contents of the three sedimentary whole-rock powders were determined on fused glass discs produced by the Fusion method (spectroflux 105) using the Automated Philips PW2404 X-ray fluorescence spectrometer at the University of Edinburgh. In-house rock standards were used to calibrate the machine and monitor accuracy and precision during analysis.

Trace element concentrations of the rock powders were determined on the PerkinElmer ELAN 6000 quadrupole ICP-MS at Durham University following the analytical procedure and instrument operating conditions described by Ottley et al. (2003). Multiple analyses of procedural blanks (3 per batch), in-house standards and international reference materials (W2, BHVO-1, AGV1, BE-N and BIR1) during each session (e.g. at the start, mid-way and at the end of a run) allowed for any drift in the instrument calibration to be detected. Reproducibility (internal and external) of standard values on the ELAN were better than 5% relative standard deviation.

Preparation of whole-rock powders for Sr, Nd and Hf isotope analysis was undertaken in the Arthur Holmes Isotope Geology Laboratory (AHIGL) at Durham University. The sample dissolution procedure and chemical separation of Sr, Hf and Nd from rock samples follow that presented by Dowall et al. (2003) and Handley et al. (2008a). Sr, Nd and Hf isotope ratios were determined on the AHIGL ThermoElectron Neptune MC-ICP-MS. Details of instrument operating conditions are presented in Nowell et al. (2003) and Dowall et al. (2003). Instrumental mass bias was corrected using $^{88}\text{Sr}/^{86}\text{Sr}$ of 8.375209 (the reciprocal of the $^{86}\text{Sr}/^{88}\text{Sr}$ ratio of 0.1194), $^{146}\text{Nd}/^{145}\text{Nd}$ of 2.079143 (equivalent to the more commonly used $^{146}\text{Nd}/^{144}\text{Nd}$ ratio of 0.7219) and $^{179}\text{Hf}/^{177}\text{Hf}$ of 0.7325 using an exponential law. Data quality was monitored over several analytical sessions by regular analysis of standard reference materials during each run. The reproducibility of $^{87}\text{Sr}/^{86}\text{Sr}$ for NBS 987 in each of the analytical sessions was better than 21 ppm. The reproducibility of $^{143}\text{Nd}/^{144}\text{Nd}$ and $^{176}\text{Hf}/^{177}\text{Hf}$ for the respective standard solutions in each of the analytical sessions was better than 19 and 28 ppm (2σ), respectively. The average reproducibility and accuracy of Nd and Hf isotope ratios of standard solutions over the period of

study are shown in Table B.3 (Appendix B) of Handley et al. (2010). Total procedural blanks (at least 2 processed per sample batch) were analysed by ICP-MS on the PerkinElmer ELAN 6000 quadrupole at Durham University and were below 1.2 ng (typically <300 pg), 219 pg and 73 pg, respectively, for Sr, Nd and Hf. These values are insignificant considering the quantity of Sr, Nd and Hf processed from the rocks.

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